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(71) Applicants (for all designated States except US): THE REGENTS OF THE UNIVERSITY OF MICHIGAN [US/US]; Technology Management Office, Wolverine Tower, Room 2071, 3003 South State Street, Ann Arbor, MI 48109-1280 (US). BOARD OF TRUSTEES OPERATING MICHIGAN STATE UNIVERSITY [US/US]; East Lansing, MI 48824 (US).			
(72) Inventors; and			
(75) Inventors/Applicants (for US only): VENTA, Patrick, J. [US/US]; 9646 Rolling Green, Pinckney, MI 48169 (US). BREWER, George, J. [US/US]; 3820 Gensley, Ann Arbor, MI 48103 (US). YUZBASIYAN-GURKAN, Vilma [US/US]; 3101 Dexter Road, Ann Arbor, MI 48103 (US). SCHALL, William, D. [US/US]; 3150 S. Williamston, Williamston, MI 48895 (US).			
(74) Agents: SMITH, DeAnn, F. et al.; Harness, Dickey & Pierce, P.L.C., P.O. Box 828, Bloomfield Hills, MI 48303 (US).			
(54) Title: DNA ENCODING CANINE VON WILLEBRAND FACTOR AND METHODS OF USE			
(57) Abstract The complete sequence of the canine von Willebrand Factor cDNA and deduced amino acid sequence is provided. The mutation which causes von Willebrand's Disease in Scottish Terriers, a single base deletion in exon 4, has also been determined. Methods for detecting carriers of the defective vWF gene are also provided.			

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**DNA ENCODING CANINE VON WILLEBRAND FACTOR
AND METHODS OF USE**

FIELD OF THE INVENTION

This invention relates generally to canine von Willebrand factor (vWF), and
5 more particularly, to the gene encoding vWF as well as a genetic defect that causes
canine von Willebrand's disease.

BIOLOGICAL DEPOSITS

SEQUENCE

ACCESSION NO.

Canine von Willebrand Factor

10

BACKGROUND OF THE INVENTION

In both dogs and humans, von Willebrand's disease (vWD) is a bleeding disorder of variable severity that results from a quantitative or qualitative defect in von Willebrand factor (vWF) (Ginsburg, D. et al., *Blood* 79:2507-2519 (1992); Ruggeri, Z.M., et al., *FASEB J* 7:308-316 (1993); Dodds, W.J., *Mod Vet Pract* 681-15 686 (1984); Johnson, G.S. et al., *JAVMA* 176:1261-1263 (1988); Brooks, M., *Probl In Vet Med* 4:636-646 (1992)). This clotting factor has two known functions, stabilization of Factor VIII (hemophilic factor A) in the blood, and aiding the adhesion of platelets to the subendothelium, which allows them to provide hemostasis more effectively. If the factor is missing or defective, the patient, whether human or dog, 20 may bleed severely.

The disease is the most common hereditary bleeding disorder in both species, and is genetically and clinically heterogeneous. Three clinical types, called 1, 2, and 3 (formerly I, II, and III; see Sadler, J.E. et al., *Blood* 84:676-679 (1994) for nomenclature changes), have been described. Type 1 vWD is inherited in a 25 dominant, incompletely penetrant fashion. Bleeding appears to be due to the reduced level of vWF rather than a qualitative difference. Although this is the most common form of vWD found in most mammals, and can cause serious bleeding problems, it is generally less severe than the other two types. In addition, a relatively inexpensive vasopressin analog (DDAVP) can help alleviate symptoms 30 (Kraus, K.H. et al., *Vet Surg* 18:103-109 (1989)).

In Type 2 vWD, patients have essentially normal levels of vWF, but the factor is abnormal as determined by specialized tests (Ruggeri, Z.M., et al., *FASEB J* 7:308-316 (1993); Brooks, M., *Probl In Vet Med* 4:636-646 (1992)). This type is also

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inherited in a dominant fashion and has only rarely been described in dogs (Turrentine, M.A., et al., *Vet Clin North Am Small Anim Pract* 18:275 (1988)).

5 Type 3 vWD is the most severe form of the disease. It is inherited as an autosomal recessive trait, and affected individuals have no detectable vWF in their blood. Serious bleeding episodes require transfusions of blood or cryoprecipitate to supply the missing vWF. Heterozygous carriers have moderately reduced factor concentrations, but generally appear to have normal hemostasis.

10 Scottish terriers have Type 3 vWD (Dodds, W.J., *Mod Vet Pract* 681-686 (1984); Johnson, G.S. et al., *JAVMA* 176:1261-1263 (1988)). Homozygotes have no detectable vWF and have a severe bleeding disorder. Heterozygotes have reduced levels of the factor, and are clinically normal (Brooks, M. et al., *JAVMA* 200:1123-1127 (1992)). The prevalence of vWD among Scottish terriers including both heterozygotes and homozygotes has been variously estimated from 27-31% (Stokol, T. et al., *Res. Vet. Sci.* 59:152-155 (1995); Brooks, M., *Proc. 9th ACVIM* 15 *Forum* 89-91 (1991)).

20 Currently, detection of affected and carrier Scottish terrier dogs is done by vWF antigen testing (Benson, R.E. et al., *Am J Vet Res* 44:399-403 (1983); Stokol, T. et al., *Res. Vet. Sci.* 59:152-155 (1995)) or by coagulation assays (Rosborough, T.K. et al., *J. Lab. Clin. Med.* 96:47-56 (1980); Read, M.S. et al., *J. Lab. Clin. Med.* 101:74-82 (1983)). These procedures yield variable results, as the protein-based tests can be influenced by such things as sample collection, sample handling, estrous, pregnancy, vaccination, age, and hypothyroidism (Strauss, H.S. et al., *New Eng J Med* 269:1251-1252 (1963); Bloom, A.L., *Mayo Clin Proc* 66:743-751 (1991); Stirling, Y. et al., *Thromb Haemostasis* 52:176-182 (1984); Mansell, P.D. et al., *Br. Vet. J.* 148:329-337 (1992); Avgeris, S. et al., *JAVMA* 196:921-924 (1990); Panciera, D.P. et al., *JAVMA* 205:1550-1553 (1994)). Thus, for example, a dog that tests within the normal range on one day, can test within the carrier range on another day. It is therefore difficult for breeders to use this information.

30 It would thus be desirable to provide the nucleic acid sequence encoding canine vWF. It would also be desirable to provide the genetic defect responsible for canine vWD. It would further be desirable to obtain the amino acid sequence of canine vWF. It would also be desirable to provide a method for detecting carriers of the defective vWF gene based on the nucleic acid sequence of the normal and defective vWF gene.

SUMMARY OF THE INVENTION

The present invention provides a novel purified and isolated nucleic acid sequence encoding canine vWF. A nucleic acid sequence containing the mutation that causes vWD in Scottish terriers, a single-base deletion in exon 4, is also provided. The nucleic acid sequences of the present invention may be used in methods for detecting carriers of the mutation that causes vWD. Such methods may be used by breeders to reduce the frequency of the disease-causing allele and the incidence of disease. In addition, the nucleic acid sequence of the canine vWF provided herein may be used to determine the genetic defect that causes vWD in other breeds as well as other species.

Additional objects, advantages, and features of the present invention will become apparent from the following description, taken in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

The various advantages of the present invention will become apparent to one skilled in the art by reading the following specification and by referencing the following drawings in which:

Figures 1A-1C is the nucleic acid sequence of the canine von Willebrand factor of the present invention;

Figures 2A-2C is a comparison of the human and canine prepro-von Willebrand factor amino acid sequences;

Figure 3 provides nucleotide sequencing ladders for the von Willebrand's disease mutation region for normal (clear), carrier, and affected Scottish terriers, the sequences being obtained directly from PCR products derived from genomic DNAs in exon 4;

Figure 4 illustrates the results of a method of the present invention used to detect the Scottish terrier vWD mutation; and

Figure 5 shows the Scottish terrier pedigree, which in turn illustrates segregation of the mutant and normal vWF alleles.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The cDNA encoding canine von Willebrand Factor (vWF) has been sequenced, and its sequence is set forth in Figures 1A-1C and SEQ ID NO: 1. The amino acid sequence corresponding to the cDNA of canine vWF has been subsequently deduced and is set forth in Figures 2A-2C and SEQ ID NO: 2. The mutation of the normal vWF gene which causes von Willebrand's Disease (vWD),

a deletion at codon 88 of the normal gene resulting in a frameshift, is also provided. The nucleic acid sequences of the present invention may be used in methods for detecting homozygous and heterozygous carriers of the defective vWF gene.

In a preferred method of detecting the presence of the von Willebrand allele 5 in canines, DNA samples are first collected by relatively noninvasive techniques, i.e., DNA samples are obtained with minimal penetration into body tissues of the animals to be tested. Common noninvasive tissue sample collection methods may be used and include withdrawing buccal cells via cheek swabs and withdrawing blood samples. Following isolation of the DNA by standard techniques, PCR is performed 10 on the DNA utilizing pre-designed primers that produce enzyme restriction sites on those DNA samples that harbor the defective gene. Treatment of the amplified DNA with appropriate restriction enzymes such as *Bsi*E I thus allows one to analyze for the presence of the defective allele. One skilled in the art will appreciate that this method may be applied not only to Scottish terriers, but to other breeds such as 15 Shetland sheepdogs and Dutch Kooikers.

Overall, the present invention provides breeders with an accurate, definitive test whereby the undesired vWD gene may be eliminated from breeding lines. The current tests used by breeders are protein- based, and as noted previously, the primary difficulty with this type of test is the variability of results due to a variety of 20 factors. The ultimate result of such variability is that an inordinate number of animals fall into an ambiguous grouping whereby carriers and noncarriers cannot be reliably distinguished. The present invention obviates the inherent limitations of protein-based tests by detecting the genetic mutation which causes vWD. As described in Specific Example 1, the methods of the present invention provide an 25 accurate test for distinguishing noncarriers, homozygous carriers and heterozygous carriers of the defective vWF gene.

It will be appreciated that because the vWF cDNA of the present invention is substantially homologous to vWF cDNA throughout the canine species, the nucleic acid sequences of the present invention may be used to detect DNA mutations in 30 other breeds as well. In addition, the canine vWF sequence presented herein potentially in combination with the established human sequence (Genbank Accession No. X04385, Bonthron, D. et al., *Nucleic Acids Res.* 14:7125-7128 (1986); Mancuso, D.J. et al., *Biochemistry* 30:253-269 (1989); Meyer, D. et al., *Throm Haemostasis* 70:99-104 (1993)), may be used to facilitate sequencing of the vWF

gene and genetic defects causing vWD, in other mammalian species e.g., by using cross-species PCR methods known by those skilled in the art.

It is also within the contemplation of this invention that the isolated and purified nucleic acid sequences of the present invention be incorporated into an appropriate recombinant expression vector, e.g., viral or plasmid, which is capable of transforming an appropriate host cell, either eukaryotic (e.g., mammalian) or prokaryotic (e.g., *E. coli*). Such DNA may involve alternate nucleic acid forms, such as cDNA, gDNA, and DNA prepared by partial or total chemical synthesis. The DNA may also be accompanied by additional regulatory elements, such as promoters, operators and regulators, which are necessary and/or may enhance the expression of the vWF gene product. In this way, cells may be induced to over-express the vWF gene, thereby generating desired amounts of the target vWF protein. It is further contemplated that the canine vWF polypeptide sequence of the present invention may be utilized to manufacture canine vWF using standard synthetic methods. One skilled in the art will also note that the defective protein encoded by the defective vWF gene of the present invention may also be of use in formulating a complementary diagnostic test for canine vWD that may provide further data in establishing the presence of the defective allele. Thus, production of the defective vWF polypeptide, either through expression in transformed host cells as described above for the active vWF polypeptide or through chemical synthesis, is also contemplated by the present invention.

The term "gene" as to referred herein means a nucleic acid which encodes a protein product. The term "nucleic acid" refers to a linear array of nucleotides and nucleosides, such as genomic DNA, cDNA and DNA prepared by partial or total chemical synthesis from nucleotides. The term "encoding" means that the nucleic acid may be transcribed and translated into the desired polypeptide. "Polypeptide" refers to amino acid sequences which comprise both full-length proteins and fragments thereof. "Mutation" as referred to herein includes any alteration in a nucleic acid sequence including, but not limited to, deletions, substitutions and additions.

As referred to herein, the term "capable of hybridizing under high stringency conditions" means annealing a strand of DNA complementary to the DNA of interest under highly stringent conditions. Likewise, "capable of hybridizing under low stringency conditions" refers to annealing a strand of DNA complementary to the DNA of interest under low stringency conditions. In the present invention, hybridizing

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under either high or low stringency conditions would involve hybridizing a nucleic acid sequence (e.g., the complementary sequence to SEQ ID NO: 1 or portion thereof), with a second target nucleic acid sequence. "High stringency conditions" for the annealing process may involve, for example, high temperature and/or low salt 5 content, which disfavor hydrogen bonding contacts among mismatched base pairs. "Low stringency conditions" would involve lower temperature, and/or lower salt concentration than that of high stringency conditions. Such conditions allow for two DNA strands to anneal if substantial, though not near complete complementarity exists between the two strands, as is the case among DNA strands that code for the 10 same protein but differ in sequence due to the degeneracy of the genetic code. Appropriate stringency conditions which promote DNA hybridization, for example, 6X SSC at about 45 °C, followed by a wash of 2X SSC at 50 °C are known to those skilled in the art or can be found in *Current Protocols in Molecular Biology*, John Wiley & Sons, NY (1989), 6.31-6.3.6. For example, the salt concentration in the 15 wash step can be selected from a low stringency of about 2X SSC at 50 °C to a high stringency of about 0.2X SSC at 50 °C. In addition, the temperature in the wash step can be increased from low stringency at room temperature, about 22 °C, to high stringency conditions, at about 65 °C. Other stringency parameters are described in Maniatis, T., et al., *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor 20 Laboratory Press, Cold Spring NY, (1982), at pp. 387-389; see also Sambrook, J. et al., *Molecular Cloning: A Laboratory Manual*, Second Edition, Volume 2, Cold Spring Harbor Laboratory Press, Cold Spring, NY at pp. 8.46-8.47 (1989).

SPECIFIC EXAMPLE 1

Materials And Methods

25 **Isolation of RNA.** The source of the RNA was a uterus from a Scottish Terrier affected with vWD (factor level < 0.1% and a clinical bleeder), that was surgically removed because of infection. Spleen tissue was obtained from a Doberman Pinscher affected with vWD that died from dilated cardiomyopathy (factor level 7% and a clinical bleeder). Total RNA was extracted from the tissues using 30 Trizol (Life Technologies, Gaithersburg, MD). The integrity of the RNA was assessed by agarose gel electrophoresis.

35 **Design of PCR primer sets.** Primers were designed to a few regions of the gene, where sequences from two species were available (Lavergne, J.M. et al., *Biochem Biophys Res Commun* 194:1019-1024 (1993); Bakhshi, M.R. et al., *Biochem Biophys Acta* 1132:325-328 (1992)). These primers were designed using

rules for cross-species' amplifications (Venta et al., "Genes-Specific Universal Mammalian Sequence-Tagged Sites: Application To The Canine Genome" *Biochem. Genet.* (1996) in press). Most of the primers had to be designed to other regions of the gene using the human sequence alone (Mancuso, D.J. et al., *Biochemistry* 5 30:253-269 (1991)). Good amplification conditions were determined by using human and canine genomic DNAs.

Reverse Transcriptase-PCR. Total RNA was reverse transcribed using random primers (Bergenhem, N.C.H. et al., *PNAS (USA)* 89:8789-8802 (1992)). The cDNA was amplified using the primer sets shown to work on canine genomic DNA.

10 **DNA Sequence Analysis.** Amplification products of the predicted sizes were isolated from agarose gels by adsorption onto silica gel particles using the manufacturer's method (Qiagen, Chatsworth, CA). Sequences were determined using ³²P-5' end-labeled primers and a cycle sequencing kit (United States Biochemical Corp., Cleveland, OH). The sequences of the 5' and 3' untranslated 15 regions were determined after amplification using Marathon™ RACE kits (Clontech, Palo Alto, CA). Sequences were aligned using the Eugene software analysis package (Lark Technologies, Houston, TX). The sequence of the canine intron four was determined from PCR-amplified genomic DNA.

12 **Design of a Diagnostic Test.** PCR mutagenesis was used to create 20 diagnostic and control *BsiE* I and *Sau96* I restriction enzyme sites for the test. Amplification conditions for the test are: 94°C, 1 min, 61°C, 1 min, and 72°C, 1 min, for 50 cycles using cheek swab DNA (Richards, B. et al., *Human Molecular Genetics* 2:159-163 (1992)).

16 **Population Survey.** DNA was collected from 87 Scottish terriers from 25 pedigrees. DNA was isolated either from blood using standard procedures (Sambrook, J. et al., Cold Harbor Spring Lab, Cold Harbor Spring NY, 2nd Edition, (1989)) or by cheek swab samples (Richards, B. et al., *Human Molecular Genetics* 2:159-163 (1992)). The genetic status of each animal in the survey was determined 30 using the *BsiE* I test described above.

Results

32 **Comparison of the canine and human sequences.** The alignment of the canine and human prepro-von Willebrand Factor amino acid sequences is shown in Figures 2A-2C. The location of the Scottish terrier vWD mutation is indicated by the ***. Potential N-glycosylation sites are shown in bold type. The known and 35 postulated integrin binding sites are boxed. Amino acid numbers are shown on the

right side of the figure. The human sequence is derived from Genbank accession number X04385 (Bontron, D. et al., *Nucleic Acids Res.* 14:7125-7128 (1986)).

Overall, 85.1% sequence identity is seen between the prepro-vWF sequences. The pro-region is slightly less conserved than the mature protein (81.4% 5 vs. 87.5%). There were no other noteworthy percentage sequence identity differences seen in other regions of the gene, or between the known repeats contained within the gene (data not shown). Fourteen potential N-linked glycosylation sites are present in the canine sequence, all of which correspond to similar sites contained within the human sequence. The two integrin binding sites 10 identified in the human vWF protein sequence (Lankhof, H. et al., *Blood* 86:1035-1042 (1995)) are conserved in the canine sequence as well (Figures 2A-2C). The 5' and 3' untranslated regions have diverged to a greater extent than the coding region (data not shown), comparable to that found between the human and bovine sequences derived for the 5' flanking region (Janel, N. et al., *Gene* 167:291-295 15 (1995)). Additional insights into the structure and function of the von Willebrand factor can be gained by comparison of the complete human sequence (Mancuso, D.J. et al., *Biochemistry* 30:253-269 (1989); Meyer, D. et al., *Throm Haemostasis* 70:99-104 (1993)) and the complete canine sequence reported here.

The sequence for most of exon 28 was determined (Mancuso, D.J. et al., 20 *Thromb Haemost* 69:980 (1993); Porter, C.A. et al., *Mol Phylogenet Evol* 5:89-101 (1996)). All three sequences are in complete agreement, although two silent variants have been found in other breeds (Table 1, exon 28). Partial sequences of 25 exons 40 and 41 (cDNA nucleotide numbers 6923 to 7155, from the initiation codon) were also determined as part of the development of a polymorphic simple tandem repeat genetic marker (Shibuya, H. et al., *Anim Genet* 24:122 (1994)). There is a single nucleotide sequence difference between this sequence ("T") and the sequence of the present invention, ("C") at nucleotide position 6928.

Scottish Terrier vWD mutation. Figure 3 shows nucleotide sequencing ladders for the von Willebrand's Disease mutation region for normal (clear), carrier, 30 and affected Scottish terriers. The sequences were obtained directly from PCR products derived from genomic DNAs in exon 4. The arrowheads show the location of the C nucleotide that is deleted in the disease-causing allele. Note that in the carrier ladder each base above the point of the mutation has a doublet appearance, as predicted for deletion mutations. The factor levels reported for these animals 35 were: Normal, 54%; Carrier, 34%; Affected, <0.1%.

As a result of the deletion, a frameshift mutation at codon 88 leads to a new stop codon 103 bases downstream. The resulting severely truncated protein of 119 amino acids does not include any of the mature von Willebrand factor region. The identity of the base in the normal allele was determined from an unaffected dog.

5 *Development of a diagnostic test.* A PCR primer was designed to produce a *BsiE* I site in the mutant allele but not in the normal allele (Figure 4). The position of the deleted nucleotide is indicated by an asterisk. The altered nucleotides in each primer are underlined. The normal and mutant allele can also be distinguished using *Sau96* I. The naturally occurring *Sau96* I sites are shown by double underlines.

10 The highly conserved donor and acceptor dinucleotide splice sequences are shown in bold type.

In order to ensure that the restriction enzyme cut the amplified DNA to completion, an internal control restriction site common to both alleles was designed into the non-diagnostic primer. The test was verified by digestion of the DNA from 15 animals that were affected, obligate carriers, or normal (based on high factor levels [greater than 100% of normal] obtained from commonly used testing labs and reported to us by the owners, and also using breeds in which Type 3 vWD has not been observed). The expected results were obtained (e.g., Figure 5). Five vWD-affected animals from a colony founded from Scottish terriers (Brinkhous, K.M. et al., 20 *Ann. New York Acad. Sci.* 370:191-203 (1981)) were also shown to be homozygous for this mutation. An additional unaffected animal from this same colony was found to be clear.

It would still be possible to misinterpret the results of the test if restriction enzyme digestion was not complete, and if the rates of cleavage of the cont778rol 25 and diagnostic sites were vastly different. The rates of cleavage of the two *BsiE* I sites were thus examined by partially digesting the PCR products and running them on capillary electrophoresis. The rates were found to be very nearly equal (the diagnostic site is cut 12% faster than the control site).

The mutagenesis primer was also designed to produce a *Sau96* I site into the 30 normal allele but not the mutant allele. This is the reverse relationship compared to the *BsiE* I-dependent test, with respect to which allele is cut. Natural internal *Sau96* I sites serve as digestion control sites (shown in Figure 4). The test using this enzyme produced identical genotypic results compared to the *BsiE* I for all animals examined (data not shown).

- 10 -

A possible mutation in the Doberman Pinscher gene. The complete Scottish terrier sequence was compared to the complete Doberman Pinscher sequence. Several nucleotide differences were found and were compared to the nucleotides found in the same position in the human sequence as shown in Table 5 below. Most of these changes were silent. However, of three amino acid changes, one is relatively non-conservative (F905L) and is proposed to be the mutation that causes Doberman Pinscher vWD. Other data strongly suggest that the nucleotide interchange at the end of exon 43 causes a cryptic splice site to be activated reducing the amount of normally processed mRNA, with a concomitant 10 decrease in the amount of vWF produced.

Mendelian inheritance. One test often used to verify the correct identification of a mutant allele is its inheritance according to Mendel's law of segregation. Three pedigrees were examined in which the normal and mutant alleles were segregating, as shown in Figure 5. Exon four of the vWF gene was 15 PCR-amplified from genomic DNA. The PCR products were examined for the presence of the normal and mutant vWF alleles by agarose gel electrophoresis after digestion with *Bs*I (see Figure 5). The affected animals are homozygous for the mutant allele (229 bp; lanes 3 and 5). The other animals in this pedigree are heterozygotes (251 bp and 229 bp; lanes 1, 2, 4, and 6), including the obligate 20 carrier parents.

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Table 1 - Differences Between Scottie And Doberman
Protein And Nucleotide von Willebrand Factor Sequences
With Comparison To The Human Sequences

Exon	A.A. ¹	Amino Acid			Codon		
		Human	Scottie	Doberman	Human	Scottie	Doberman
5	5' UT ²	nuc - 35 ³	N/A ⁴	N/A	N/A	A	G
	4	85	S	S/F.Shift ⁵	S	TCC	TCC/TC
	5	173	M	R	K	ATG	AGG
	11	422	S	T	T	TCC	ACA
	21	898	C	C	G	TGC	TGT
10	21	905	F	F	L	TTT	TTC
	24	1041	S	S	S	TCA	TCA
	24	1042	S	S	S	TCC	TCC
	28	1333	D	D	E	GAC	GAC
	28	1349	Y	Y	Y	TAT	TAT
15	42	2381	P	L	P	CCC	CTG
	43	2479	S	S	S	TCG	TCG
	45	2555	P	P	P	CCC	CCC
	47	2591	P	P	P	CCC	CCT
	49	2672	D	D	D	GAT	GAT
20	51	2744	E	E	E	GAG	GAC
							GAA

¹Amino acid residue position²Untranslated region³Nucleotide position⁴Not Applicable⁵Frameshift mutation

Boxed residues show amino acid differences between breeds

*This site has been shown to be polymorphic in some breeds

The mature VWF protein begins in exon 18

The alleles, as typed by both the *Bs*I and *Sau*96 I tests, showed no
30 inconsistencies with Mendelian inheritance. One of these pedigrees included two
affected animals, two phenotypically normal siblings, and the obligate carrier parents.
The two parents were found to be heterozygous by the test, the two affected animals
were found to be homozygous for the mutant allele, and the normal siblings were
found to be heterozygotes.

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Population survey for the mutation: Cheek swabs or blood samples were collected from 87 animals in order to determine the incidence of carriers in the U.S. Scottish terrier population. Although we attempted to make the sample as random as possible, these dogs were found to come from 16 pedigrees, several of which are 5 more distantly interconnected. This is due to some ascertainment bias, based on ownership (as opposed to phenotypic ascertainment bias). In these 87 animals four affected and 15 carrier animals were found.

Discussion

These results establish that the single base deletion found in exon four of the 10 vWF gene causes vWD in the Scottish terrier breed. The protein produced from the mutant allele is extremely short and does not include any of the mature vWF protein. Four Scottish terriers known to be affected with the disease are homozygous for the mutation. Five other mixed-breed dogs descended from Scottish terriers, and 15 affected with vWD, are also homozygous for the mutation. No normal animals are homozygous for the mutation. Unaffected obligate carriers are always heterozygous for the mutation.

The gene frequency, as determined from the population survey, appears to be around 0.13 resulting in a heterozygote frequency of about 23% and expected 20 frequency of affected animals of about 2%. Although the sample size is relatively small and somewhat biased, these data are in general agreement with the protein-based surveys (Stokol, T. et al., *Res Vet Sci* 59:152-155 (1995); Brooks, M., *Probl In Vet Med* 4:636-646 (1992)), in that the allele frequency is substantial.

All data collected thus far indicate that this mutation accounts for essentially 25 all of the von Willebrand's disease found in Scottish terriers. This result is consistent with the results found for other genetic diseases, defined at the molecular level, in various domestic animals (Shuster, D.E. et al., *PNAS (USA)* 89:9225-9229 (1992); Rudolph, J.A. et al., *Nat Genet* 2:144-147 (1992); O'Brien, P.J. et al., *JAVMA* 203:842-851 (1993)). A likely explanation may be found in the pronounced founder 30 effect that occurs in domestic animals, compared to most human and wild animal populations.

Published data using the protein-based factor assays have shown that, at 35 least in several instances, obligate carriers have had factor levels that would lead to a diagnosis of "clear" of the disease allele. For example, in one study an obligate carrier had a factor level of 78% (Johnson, G.S. et al., *JAVMA* 176:1261-1263 (1980)). In another study, at least some of the obligate carriers had factor levels of

65% or greater (Brinkhous, K.M. et al., *Ann. New York Acad. Sci.* 370:191-203 (1981)). In addition, the number of animals that fall into an equivocal range can be substantial. In one study, 19% of Scottish terriers fell in this range (50-65% of the normal vWF antigen level) (Stokol, T. et al., *Res Vet Sci* 59:152-155 (1995)). Thus, 5 although the protein-based tests have been useful, the certainty of the DNA-based test described herein should relieve the necessity of repeated testing and the variability associated with the protein-based assays.

The mutation is present in the pre-vWF part of the molecule. This part of the molecule is processed off prior to delivery of the mature protein into the plasma. 10 This pre-portion of the molecule is important for the assembly of the mature vWF protein (Verwiej, L. et al., *EBMO J* 6:2885-2890 (1987); Wise, R.J. et al., *Cell* 52:229-236 (1988)). With the Scottish terrier frameshift vWD mutation, neither this pre-portion nor any of the mature factor is ever produced, in keeping with the fact that no factor has ever been detected in the blood of affected dogs.

15 The determination of the complete canine vWF cDNA sequence will have an impact upon the development of carrier tests for other breeds and other species as well. Currently, Shetland sheepdogs and Dutch Kooikers are known to have a significant amount of Type 3 vWD (Brooks, M. et al., *JAVMA* 200:1123-1127 (1992); Slappendel, R.J., *Vet-Q* 17:S21-S22 (1995)). Type 3 vWD has occasionally been seen 20 in other breeds as well (e.g., Johnson, G.S. et al., *JAVMA* 176:1261-1263 (1980)). All Type 3 vWD mutations described in humans to date have been found within the vWF gene itself. The availability of the canine sequence will make it easier to find the mutations in these breeds. In addition, at least some Type 1 mutations have been found within the human vWF gene, and thus Type 1 mutations may also be 25 found within the vWF gene for breeds affected with that form of the disease. The availability of two divergent mammalian vWF cDNA sequences will also make it much easier to sequence the gene from other mammalian species using cross-species PCR methods (e.g., Venta et al., *Biochem. Genet.* (1996) in press).

The test described herein for the detection of the mutation in Scottish terriers 30 may be performed on small amounts of DNA from any tissue. The tissues that are the least invasive to obtain are blood and buccal cells. For maximum convenience, a cheek swab as a source of DNA is preferred.

The foregoing discussion discloses and describes merely exemplary 35 embodiments of the present invention. One skilled in the art will readily recognize from such discussion, and from the accompanying drawings, that various changes,

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modifications and variations can be made therein without departing from the spirit and scope of the invention.

All patents and other publications cited herein are expressly incorporated by reference.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Venta, Patrick J
Yuzbasiyan-Gurkan, Vilma
Schall, William D
Brewer, George J
- (ii) TITLE OF INVENTION: DNA ENCODING CANINE VON WILLEBRAND FACTOR AND METHODS OF USE
- (iii) NUMBER OF SEQUENCES: 2
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Harness, Dickey & Pierce, P.L.C.
 - (B) STREET: 5445 Corporate Drive
 - (C) CITY: Troy
 - (D) STATE: Michigan
 - (E) COUNTRY: USA
 - (F) ZIP: 48098
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Smith, DeAnn F.
 - (C) REFERENCE/DOCKET NUMBER: 211501226PCA
- (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 248-641-1600
 - (B) TELEFAX: 248-641-0270
 - (C) TELEX: 287637

(2) INFORMATION FOR SEQ ID NO:1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8802 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 203..8641
 - (D) OTHER INFORMATION: /function= "Blood Clotting Protein"
/product= "Canine von Willebrand Factor"
/standard_name= "vWF"

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(x) PUBLICATION INFORMATION:

(A) AUTHORS: Venta, Patrick J.
 Li, Jianping
 Yuzbasiyan-Gurkan, Vilma
 Schall, William D.
 Brewer, George J.

(B) TITLE: Von Willebrand's Disease in the Scottish
 Terrier is Caused by a Single Base Deletion in
 Exon Four of the von Willebrand Factor Gene

(C) JOURNAL: Journal of the American Veterinary Medicine Association

(G) DATE: 1996

(K) RELEVANT RESIDUES IN SEQ ID NO:1: FROM 1 TO 8802

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

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ACTTGACACA	CGGACAGTAG	TACATACCAAG	TAGCTCTCTG	CGAGGACGGT	GATCACTAAT	180
CATTTCTCCT	GCTTCGTGGC	AG ATG AGT CCT ACC AGA CTT GTG AGG GTG CTG				232
		Met Ser Pro Thr Arg Leu Val Arg Val Leu				
		1 5 10				
CTG GCT CTG	GCC CTC ATC	TTG CCA GGG AAA CTT TGT ACA AAA GGG ACT				280
Leu Ala	Leu Ala	Leu Ile Leu Pro Gly Lys Leu Cys Thr Lys Gly Thr				
15	20	25				
GTT GGA AGG	TCA TCG ATG	GCC CGA TGT AGC CTT CTC GGA GGT GAC TTC				328
Val Gly Arg	Ser Ser Met	Ala Arg Cys Ser Leu Leu Gly Gly Asp Phe				
30	35	40				
ATC AAC ACC	TTT GAT GAG	AGC ATG TAC AGC TTT GCG GGA GAT TGC AGT				376
Ile Asn Thr	Phe Asp Glu	Ser Met Tyr Ser Phe Ala Gly Asp Cys Ser				
45	50	55				
TAC CTC CTG	GCT GGG GAC	TGC CAG GAA CAC TCC ATC TCA CTT ATC GGG				424
Tyr Leu Ala	Gly Asp Cys	Gln Glu His Ser Ile Ser Leu Ile Gly				
60	65	70				
GGT TTC CAA AAT	GAC AAA AGA	GTG AGC CTC TCC GTG TAT CTC GGA GAA				472
Gly Phe Gln Asn	Asp Lys Arg	Val Ser Leu Ser Val Tyr Leu Gly Glu				
75	80	85	90			
TTT TTC GAC ATT	CAT TTG TTT	GTC AAT GGT ACC ATG CTG CAG GGG ACC				520
Phe Phe Asp Ile	His Leu Phe Val	Asn Gly Thr Met Leu Gln Gly Thr				
95	100	105				
CAA AGC ATC	TCC ATG CCC	TAC GCC TCC AAT GGG CTG TAT CTA GAG GCC				568
Gln Ser Ile	Ser Met Pro	Tyr Ala Ser Asn Gly Leu Tyr Leu Glu Ala				
110	115	120				
GAG GCT GGC	TAC TAC AAG	CTG TCC AGT GAG GCC TAC GGC TTT GTG GCC				616
Glu Ala	Gly Tyr Tyr	Lys Leu Ser Ser Glu Ala Tyr Gly Phe Val Ala				
125	130	135				
AGA ATT GAT GGC	AAT GGC AAC	TTT CAA GTC CTG CTG TCA GAC AGA TAC				664
Arg Ile Asp Gly	Asn Gly Asn	Phe Gln Val Leu Leu Ser Asp Arg Tyr				
140	145	150				
TTC AAC AAG ACC	TGT GGG CTG	TGT GGC AAC TTT AAT ATC TTT GCT GAG				712
Phe Asn Lys Thr	Cys Gly Leu Cys	Gly Asn Phe Asn Ile Phe Ala Glu				
155	160	165	170			

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GCC CGC TGC CAC CCG CTG GTG GAC CCT GAG CCT TTT GTC GCC CTG TGT Ala Arg Cys His Pro Leu Val Asp Pro Glu Pro Phe Val Ala Leu Cys 235 240 245 250	952
GAA AGG ACT CTG TGC ACC TGT GTC CAG GGG ATG GAG TGC CCT TGT GCG Glu Arg Thr Leu Cys Thr Cys Val Gln Gly Met Glu Cys Pro Cys Ala 255 260 265	1000
GTC CTC CTG GAG TAC GCC CGG GCC TGT GCC CAG CAG GGG ATT GTC TTG Val Leu Leu Glu Tyr Ala Arg Ala Cys Ala Gln Gly Ile Val Leu 270 275 280	1048
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CTT CAT GTC AAA GAA GTG TGT CAG GAG CAA TGT GTA GAT GGC TGC AGC Leu His Val Lys Glu Val Cys Gln Glu Gln Cys Val Asp Gly Cys Ser 315 320 325 330	1192
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TGC AGC AAT GAA GAA TGC CCA GGC GAG TGT CTG GTC ACA GGA CAG TCC Cys Ser Asn Glu Glu Cys Pro Gly Glu Cys Leu Val Thr Gly Gln Ser 380 385 390	1384
CAC TTC AAG AGC TTC GAC AAC AGG TAC TTC ACC TTC AGT GGG GTC TGC His Phe Lys Ser Phe Asp Asn Arg Tyr Phe Thr Phe Ser Gly Val Cys 395 400 405 410	1432
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GGA GAT TGT GTG CCC AAG GCT CAG TGT CCC TGT TAC TAT GAT GGT GAG Gly Asp Cys Val Pro Lys Ala Gln Cys Pro Cys Tyr Tyr Asp Gly Glu 700 705 710	2344

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TGT GAG GAT GGC TTC ATG CAC TGT ACC ACA AGT GGA GGC CTG GGA AGC Cys Glu Asp Gly Phe Met His Cys Thr Thr Ser Gly Gly Leu Gly Ser 735 740 745	2440
CTG CTG CCC AAC CCG GTG CTC AGC AGC CCC CGG TGT CAC CGC AGC AAA Leu Leu Pro Asn Pro Val Leu Ser Ser Pro Arg Cys His Arg Ser Lys 750 755 760	2488
AGG AGC CTG TCC TGT CGG CCC CCC ATG GTC AAG TTG GTG TGT CCC GCT Arg Ser Leu Ser Cys Arg Pro Pro Met Val Lys Leu Val Cys Pro Ala 765 770 775	2536
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CCG CAG GGC ATG CGG CAT GAA AAC AGG TGT GTG GCG CTG GAA AGA Pro Gln Gly Met Val Arg His Glu Asn Arg Cys Val Ala Leu Glu Arg 815 820 825	2680
TGT CCC TGC TTC CAC CAA GGC CAA GAG TAC GCC CCA GGA GAA ACC GTG Cys Pro Cys Phe His Gln Gly Gln Glu Tyr Ala Pro Gly Glu Thr Val 830 835 840	2728
AAA ATT GAC TGC AAC ACT TGT GTC TGT CGG GAC CGG AAG TGG ACC TGC Lys Ile Asp Cys Asn Thr Cys Val Cys Arg Asp Arg Lys Trp Thr Cys 845 850 855	2776
ACA GAC CAT GTG TGT GAT GCC ACT TGC TCT GCC ATC GGC ATG GCG CAC Thr Asp His Val Cys Asp Ala Thr Cys Ser Ala Ile Gly Met Ala His 860 865 870	2824
TAC CTC ACC TTC GAC GGA CTC AAG TAC CTG TTC CCT GGG GAG TGC CAG Tyr Leu Thr Phe Asp Gly Leu Lys Tyr Leu Phe Pro Gly Glu Cys Gln 875 880 885 890	2872
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ATC CTG GTG GGG AAC GAG GGG TGC AGC TAC CCC TCA GTG AAA TGC AAG Ile Leu Val Gly Asn Glu Gly Cys Ser Tyr Pro Ser Val Lys Cys Lys 910 915 920	2968
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GGG GAG GTG AAT GTG AAG AAA CCC ATG AAG GAT GAG ACT CAC TTT GAG Gly Glu Val Asn Val Lys Lys Pro Met Lys Asp Glu Thr His Phe Glu 940 945 950	3064
GTG GTA GAG TCT GGT CAG TAC GTC ATT CTG CTG GGC AAG GCA CTC Val Val Glu Ser Gly Gln Tyr Val Ile Leu Leu Leu Gly Lys Ala Leu 955 960 965 970	3112
TCT GTG GTC TGG GAC CAC CGC CTG AGC ATC TCT GTG ACC CTG AAG CGG Ser Val Val Trp Asp His Arg Leu Ser Ile Ser Val Thr Leu Lys Arg 975 980 985	3160

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CAG AAC AAT GAT TTC ACC AGC AGC CTC CAA ATA GAA GAA GAC CCT Gln Asn Asn Asp Phe Thr Ser Ser Leu Gln Ile Glu Glu Asp Pro 1005 1010 1015	3256
GTG GAC TTT GGG AAT TCC TGG AAA GTG AAC CCG CAG TGT GCC GAC ACC Val Asp Phe Gly Asn Ser Trp Lys Val Asn Pro Gln Cys Ala Asp Thr 1020 1025 1030	3304
AAG AAA GTA CCA CTG GAC TCA TCC CCT GCC GTC TGC CAC AAC AAC ATC Lys Lys Val Pro Leu Asp Ser Ser Pro Ala Val Cys His Asn Asn Ile 1035 1040 1045 1050	3352
ATG AAG CAG ACG ATG GTG GAT TCC TCC TGC AGG ATC CTC ACC AGT GAT Met Lys Gln Thr Met Val Asp Ser Ser Cys Arg Ile Leu Thr Ser Asp 1055 1060 1065	3400
ATT TTC CAG GAC TGC AAC AGG CTG GTG GAC CCT GAG CCA TTC CTG GAC Ile Phe Gln Asp Cys Asn Arg Leu Val Asp Pro Glu Pro Phe Leu Asp 1070 1075 1080	3448
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GTG GAG GAC ACG TCG GAG CCG CCC CTC CAT GAC TTC CAC TGC AGC AGG Val Glu Asp Thr Ser Glu Pro Pro Leu His Asp Phe His Cys Ser Arg 1260	1265	1270	4024	
CTT CTG GAC CTG GTT TTC CTG CTG GAT GGC TCC TCC AAG CTG TCT GAG Leu Leu Asp Leu Val Phe Leu Leu Asp Gly Ser Ser Lys Leu Ser Glu 1275	1280	1285	1290	4072
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GAG CTG CGG CGC ATC ACC AGC CAG GTG AAG TAC GCG GGC AGC GAG GTG Glu Leu Arg Arg Ile Thr Ser Gln Val Lys Tyr Ala Gly Ser Glu Val 1340	1345	1350	4264	
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GAT GAG ATT ATC AAC TAC CTC TGT GAC CTT GCC CCC GAA GCA CCT GCC Asp Glu Ile Ile Asn Tyr Leu Cys Asp Leu Ala Pro Glu Ala Pro Ala 1455	1460	1465	4600	
CCT ACT CAG CAC CCC CCA ATG GCC CAG GTC ACG GTG GGT TCG GAG CTG Pro Thr Gln His Pro Pro Met Ala Gln Val Thr Val Gly Ser Glu Leu 1470	1475	1480	4648	
TTG GGG GTT TCA TCT CCA GGA CCC AAA AGG AAC TCC ATG GTC CTG GAT Leu Gly Val Ser Ser Pro Gly Pro Lys Arg Asn Ser Met Val Leu Asp 1485	1490	1495	4696	
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CAG CAG GTG CGG GAT ATC CGA TAC CGG GGT GGC AAC AGG ACC AAC ACT Gln Gln Val Arg Asp Ile Arg Tyr Arg Gly Gly Asn Arg Thr Asn Thr 1565 1570 1575	4936
GGA CTG GCC CTG CAA TAC CTG TCC GAA CAC AGC TTC TCG GTC AGC CAG Gly Leu Ala Leu Gln Tyr Leu Ser Glu His Ser Phe Ser Val Ser Gln 1580 1585 1590	4984
GGG GAC CGG GAG CAG GTA CCT AAC CTG GTC TAC ATG GTC ACA GGA AAC Gly Asp Arg Glu Gln Val Pro Asn Leu Val Tyr Met Val Thr Gly Asn 1595 1600 1605 1610	5032
CCC GCT TCT GAT GAG ATC AAG CGG ATG CCT GGA GAC ATC CAG GTG GTG Pro Ala Ser Asp Glu Ile Lys Arg Met Pro Gly Asp Ile Gln Val Val 1615 1620 1625	5080
CCC ATC GGG GTG GGT CCA CAT GCC AAT GTG CAG GAG CTG GAG AAG ATT Pro Ile Gly Val Gly Pro His Ala Asn Val Gln Glu Leu Glu Lys Ile 1630 1635 1640	5128
GGC TGG CCC AAT GCC CCC ATC CTC ATC CAT GAC TTT GAG ATG CTC CCT Gly Trp Pro Asn Ala Pro Ile Leu Ile His Asp Phe Glu Met Leu Pro 1645 1650 1655	5176
CGA GAG GCT CCT GAT CTG GTG CTA CAG AGG TGC TGC TCT GGA GAG GGG Arg Glu Ala Pro Asp Leu Val Leu Gln Arg Cys Cys Ser Gly Glu Gly 1660 1665 1670	5224
CTG CAG ATC CCC ACC CTC TCC CCC ACC CCA GAT TGC AGC CAG CCC CTG Leu Gln Ile Pro Thr Leu Ser Pro Thr Pro Asp Cys Ser Gln Pro Leu 1675 1680 1685 1690	5272
GAT GTG GTC CTC CTC CTG GAT GGC TCT TCC AGC ATT CCA GCT TCT TAC Asp Val Val Leu Leu Asp Gly Ser Ser Ser Ile Pro Ala Ser Tyr 1695 1700 1705	5320
TTT GAT GAA ATG AAG AGC TTC ACC AAG GCT TTT ATT TCA AGA GCT AAT Phe Asp Glu Met Lys Ser Phe Thr Lys Ala Phe Ile Ser Arg Ala Asn 1710 1715 1720	5368
ATA GGG CCC CGG CTC ACT CAA GTG TCG GTG CTG CAA TAT GGA AGC ATC Ile Gly Pro Arg Leu Thr Gln Val Ser Val Leu Gln Tyr Gly Ser Ile 1725 1730 1735	5416
ACC ACT ATC GAT GTG CCT TGG AAT GTA GCC TAT GAG AAA GTC CAT TTA Thr Thr Ile Asp Val Pro Trp Asn Val Ala Tyr Glu Lys Val His Leu 1740 1745 1750	5464
CTG AGC CTT GTG GAC CTC ATG CAG CAG GAG GGA GGC CCC AGC GAA ATT Leu Ser Leu Val Asp Leu Met Gln Gln Glu Gly Gly Pro Ser Glu Ile 1755 1760 1765 1770	5512
GGG GAT GCT TTG AGC TTT GCC GTG CGA TAT GTC ACC TCA GAA GTC CAT Gly Asp Ala Leu Ser Phe Ala Val Arg Tyr Val Thr Ser Glu Val His 1775 1780 1785	5560
GGT GCC AGG CCC GGA GCC TCG AAA GCG GTG GTT ATC CTA GTC ACA GAT Gly Ala Arg Pro Gly Ala Ser Lys Ala Val Val Ile Leu Val Thr Asp 1790 1795 1800	5608

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GTC TCC GTG GAT TCA GTG GAT GCT GCA GCC GAG GCC GCC AGA TCC AAC Val Ser Val Asp Ser Val Asp Ala Ala Ala Glu Ala Ala Arg Ser Asn 1805 1810 1815	5656
CGA GTG ACA GTG TTC CCC ATT GGA ATC GGG GAT CGG TAC AGT GAG GCC Arg Val Thr Val Phe Pro Ile Gly Ile Gly Asp Arg Tyr Ser Glu Ala 1820 1825 1830	5704
CAG CTG AGC AGC TTG GCA GGC CCA AAG GCT GGC TCC AAT ATG GTA AGG Gln Leu Ser Ser Leu Ala Gly Pro Lys Ala Gly Ser Asn Met Val Arg 1835 1840 1845 1850	5752
CTC CAG CGA ATT GAA GAC CTC CCC ACC GTG GCC ACC CTG GGA AAT TCC Leu Gln Arg Ile Glu Asp Leu Pro Thr Val Ala Thr Leu Gly Asn Ser 1855 1860 1865	5800
TTC TTC CAC AAG CTG TGC TCT GGG TTT GAT AGA GTT TGC GTG GAT GAG Phe Phe His Lys Leu Cys Ser Gly Phe Asp Arg Val Cys Val Asp Glu 1870 1875 1880	5848
GAT GGG AAT GAG AAG AGG CCC GGG GAT GTC TGG ACC TTG CCA GAC CAG Asp Gly Asn Glu Lys Arg Pro Gly Asp Val Trp Thr Leu Pro Asp Gln 1885 1890 1895	5896
TGC CAC ACA GTG ACT TGC CTG CCA GAT GGC CAG ACC TTG CTG AAG AGT Cys His Thr Val Thr Cys Leu Pro Asp Gly Gln Thr Leu Leu Lys Ser 1900 1905 1910	5944
CAT CGG GTC AAC TGT GAC CGG GGG CCA AGG CCT TCG TGC CCC AAT GGC His Arg Val Asn Cys Asp Arg Gly Pro Arg Pro Ser Cys Pro Asn Gly 1915 1920 1925 1930	5992
CAG CCC CCT CTC AGG GTA GAG GAG ACC TGT GGC TGC CGC TGG ACC TGT Gln Pro Pro Leu Arg Val Glu Thr Cys Gly Cys Arg Trp Thr Cys 1935 1940 1945	6040
CCC TGT GTG TGC ATG GGC AGC TCT ACC CGG CAC ATC GTG ACC TTT GAT Pro Cys Val Cys Met Gly Ser Ser Thr Arg His Ile Val Thr Phe Asp 1950 1955 1960	6088
GGG CAG AAT TTC AAG CTG ACT GGC AGC TGT TCG TAT GTC CTA TTT CAA Gly Gln Asn Phe Lys Leu Thr Gly Ser Cys Ser Tyr Val Leu Phe Gln 1965 1970 1975	6136
AAC AAG GAG CAG GAC CTG GAG GTG ATT CTC CAG AAT GGT GCC TGC AGC Asn Lys Glu Gln Asp Leu Glu Val Ile Leu Gln Asn Gly Ala Cys Ser 1980 1985 1990	6184
CCT GGG GCG AAG GAG ACC TGC ATG AAA TCC ATT GAG GTG AAG CAT GAC Pro Gly Ala Lys Glu Thr Cys Met Lys Ser Ile Glu Val Lys His Asp 1995 2000 2005 2010	6232
GGC CTC TCA GTT GAG CTC CAC AGT GAC ATG CAG ATG ACA GTG AAT GGG Gly Leu Ser Val Glu Leu His Ser Asp Met Gln Met Thr Val Asn Gly 2015 2020 2025	6280
AGA CTA GTC TCC ATC CCA TAT GTG GGT GGA GAC ATG GAA GTC AAT GTT Arg Leu Val Ser Ile Pro Tyr Val Gly Gly Asp Met Glu Val Asn Val 2030 2035 2040	6328
TAT GGG ACC ATC ATG TAT GAG GTC AGA TTC AAC CAT CTT GGC CAC ATC Tyr Gly Thr Ile Met Tyr Glu Val Arg Phe Asn His Leu Gly His Ile 2045 2050 2055	6376
TTC ACA TTC ACC CCC CAA AAC AAT GAG TTC CAG CTG CAG CTC AGC CCC Phe Thr Phe Thr Pro Gln Asn Asn Glu Phe Gln Leu Gln Leu Ser Pro 2060 2065 2070	6424

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AGG ACC TTT GCT TCG AAG ACA TAT GGT CTC TGT GGG ATC TGT GAT GAG Arg Thr Phe Ala Ser Lys Thr Tyr Gly Leu Cys Gly Ile Cys Asp Glu 2075 2080 2085 2090	6472
AAC GGA GCC AAT GAC TTC ATT CTG AGG GAT GGG ACA GTC ACC ACA GAC Asn Gly Ala Asn Asp Phe Ile Leu Arg Asp Gly Thr Val Thr Thr Asp 2095 2100 2105	6520
TGG AAG GCA CTC ATC CAG GAA TGG ACC GTA CAG CAG CTT GGG AAG ACA Trp Lys Ala Leu Ile Gln Glu Trp Thr Val Gln Gln Leu Gly Lys Thr 2110 2115 2120	6568
TCC CAG CCT GTC CAT GAG GAG CAG TGT CCT GTC TCC GAA TTC TTC CAC Ser Gln Pro Val His Glu Glu Gln Cys Pro Val Ser Glu Phe Phe His 2125 2130 2135	6616
TGC CAG GTC CTC CTC TCA GAA TTG TTT GCC GAG TGC CAC AAG GTC CTC Cys Gln Val Leu Leu Ser Glu Leu Phe Ala Glu Cys His Lys Val Leu 2140 2145 2150	6664
GCT CCA GCC ACC TTT TAT GCC ATG TGC CAG CCC GAC AGT TGC CAC CCG Ala Pro Ala Thr Phe Tyr Ala Met Cys Gln Pro Asp Ser Cys His Pro 2155 2160 2165 2170	6712
AAG AAA GTG TGT GAG GCG ATT GCC TTG TAT GCC CAC CTC TGT CGG ACC Lys Lys Val Cys Glu Ala Ile Ala Leu Tyr Ala His Leu Cys Arg Thr 2175 2180 2185	6760
AAA GGG GTC TGT GTG GAC TGG AGG AGG GCC AAT TTC TGT GCT ATG TCA Lys Gly Val Cys Val Asp Trp Arg Arg Ala Asn Phe Cys Ala Met Ser 2190 2195 2200	6808
TGT CCA CCA TCC CTG GTG TAC AAC CAC TGT GAG CAT GGC TGC CCT CGG Cys Pro Pro Ser Leu Val Tyr Asn His Cys Glu His Gly Cys Pro Arg 2205 2210 2215	6856
CTC TGT GAA GGC AAT ACA AGC TCC TGT GGG GAC CAA CCC TCG GAA GGC Leu Cys Glu Gly Asn Thr Ser Ser Cys Gly Asp Gln Pro Ser Glu Gly 2220 2225 2230	6904
TGC TTC TGC CCC CCA AAC CAA GTC ATG CTG GAA GGT AGC TGT GTC CCC Cys Phe Cys Pro Pro Asn Gln Val Met Leu Glu Gly Ser Cys Val Pro 2235 2240 2245 2250	6952
GAG GAG GCC TGT ACC CAG TGC ATC AGC GAG GAT GGA GTC CGG CAC CAG Glu Glu Ala Cys Thr Gln Cys Ile Ser Glu Asp Gly Val Arg His Gln 2255 2260 2265	7000
TTC CTG GAA ACC TGG GTC CCA GCC CAC CAG CCT TGC CAG ATC TGC ACG Phe Leu Glu Thr Trp Val Pro Ala His Gln Pro Cys Gln Ile Cys Thr 2270 2275 2280	7048
TGC CTC AGT GGG CGG AAG GTC AAC TGT ACG TTG CAG CCC TGC CCC ACA Cys Leu Ser Gly Arg Lys Val Asn Cys Thr Leu Gln Pro Cys Pro Thr 2285 2290 2295	7096
GCC AAA GCT CCC ACC TGT GGC CCG TGT GAA GTG GCC CGC CTC CGC CAG Ala Lys Ala Pro Thr Cys Gly Pro Cys Glu Val Ala Arg Leu Arg Gln 2300 2305 2310	7144
AAC GCA GTG CAG TGC TGC CCG GAG TAC GAG TGT GTG TGT GAC CTG GTG Asn Ala Val Gln Cys Cys Pro Glu Tyr Glu Cys Val Cys Asp Leu Val 2315 2320 2325 2330	7192
AGC TGT GAC CTG CCC CCG GTG CCT CCC TGC GAA GAT GGC CTC CAG ATG Ser Cys Asp Leu Pro Pro Val Pro Pro Cys Glu Asp Gly Leu Gln Met 2335 2340 2345	7240

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ACC CTG ACC AAT CCT GGC GAG TGC AGA CCC AAC TTC ACC TGT GCC TGC Thr Leu Thr Asn Pro Gly Glu Cys Arg Pro Asn Phe Thr Cys Ala Cys 2350 2355 2360	7288
AGG AAG GAT GAA TGC AGA CGG GAG TCC CCG CCC TCT TGT CCC CCG CAC Arg Lys Asp Glu Cys Arg Arg Glu Ser Pro Pro Ser Cys Pro Pro His 2365 2370 2375	7336
CGG ACG CCG GCC CTT CGG AAG ACT CAG TGC TGT GAT GAG TAT GAG TGT Arg Thr Pro Ala Leu Arg Lys Thr Gln Cys Cys Asp Glu Tyr Glu Cys 2380 2385 2390	7384
GCA TGC AAC TGT GTC AAC TCC ACG GTG AGC TGC CCG CTT GGG TAC CTG Ala Cys Asn Cys Val Asn Ser Thr Val Ser Cys Pro Leu Gly Tyr Leu 2395 2400 2405 2410	7432
GCC TCG GCT GTC ACC AAC GAC TGT GGC TGC ACC ACA ACA ACC TGC TTC Ala Ser Ala Val Thr Asn Asp Cys Gly Cys Thr Thr Thr Cys Phe 2415 2420 2425	7480
CCT GAC AAG GTG TGT GTC CAC CGA GGC ACC ATC TAC CCT GTG GGC CAG Pro Asp Lys Val Cys Val His Arg Gly Thr Ile Tyr Pro Val Gly Gln 2430 2435 2440	7528
TTC TGG GAG GAG GCC TGT GAC GTG TGC ACC TGC ACG GAC TTG GAG GAC Phe Trp Glu Glu Ala Cys Asp Val Cys Thr Cys Thr Asp Leu Glu Asp 2445 2450 2455	7576
TCT GTG ATG GGC CTG CGT GTG GCC CAG TGC TCC CAG AAG CCC TGT GAG Ser Val Met Gly Leu Arg Val Ala Gln Cys Ser Gln Lys Pro Cys Glu 2460 2465 2470	7624
GAC AAC TGC CTG TCA GGC TTC ACT TAT GTC CTT CAT GAA GGC GAG TGC Asp Asn Cys Leu Ser Gly Phe Thr Tyr Val Leu His Glu Gly Glu Cys 2475 2480 2485 2490	7672
TGT GGA AGG TGT CTG CCA TCT GCC TGT GAG GTG GTC ACT GGT TCA CCA Cys Gly Arg Cys Leu Pro Ser Ala Cys Glu Val Val Thr Gly Ser Pro 2495 2500 2505	7720
CGG GGC GAC GCC CAG TCT CAC TGG AAG AAT GTT GGC TCT CAC TGG GCC Arg Gly Asp Ala Gln Ser His Trp Lys Asn Val Gly Ser His Trp Ala 2510 2515 2520	7768
TCC CCT GAC AAC CCC TGC CTC ATC AAT GAG TGT GTC CGA GTG AAG GAA Ser Pro Asp Asn Pro Cys Leu Ile Asn Glu Cys Val Arg Val Lys Glu 2525 2530 2535	7816
GAG GTC TTT GTG CAA CAG AGG AAT GTC TCC TGC CCC CAG CTG AAT GTC Glu Val Phe Val Gln Gln Arg Asn Val Ser Cys Pro Gln Leu Asn Val 2540 2545 2550	7864
CCC ACC TGC CCC ACG GGC TTC CAG CTG AGC TGT AAG ACC TCA GAG TGT Pro Thr Cys Pro Thr Gly Phe Gln Leu Ser Cys Lys Thr Ser Glu Cys 2555 2560 2565 2570	7912
TGT CCC ACC TGT CAC TGC GAG CCC CTG GAG GCC TGC TTG CTC AAT GGT Cys Pro Thr Cys His Cys Glu Pro Leu Glu Ala Cys Leu Leu Asn Gly 2575 2580 2585	7960
ACC ATC ATT GGG CCG GGG AAA AGT CTG ATG ATT GAT GTG TGT ACA ACC Thr Ile Ile Gly Pro Gly Lys Ser Leu Met Ile Asp Val Cys Thr Thr 2590 2595 2600	8008
TGC CGC TGC ACC GTG CCG GTG GGA GTC ATC TCT GGA TTC AAG CTG GAG Cys Arg Cys Thr Val Pro Val Gly Val Ile Ser Gly Phe Lys Leu Glu 2605 2610 2615	8056

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GGC AGG AAG ACC ACC TGT GAG GCA TGC CCC CTG GGT TAT AAG GAA GAG Gly Arg Lys Thr Thr Cys Glu Ala Cys Pro Leu Gly Tyr Lys Glu Glu 2620 2625 2630	8104
AAG AAC CAA GGT GAA TGC TGT GGG AGA TGT CTG CCT ATA GCT TGC ACC Lys Asn Gln Gly Glu Cys Cys Gly Arg Cys Leu Pro Ile Ala Cys Thr 2635 2640 2645 2650	8152
ATT CAG CTA AGA GGA GGA CAG ATC ATG ACA CTG AAG CGT GAT GAG ACT Ile Gln Leu Arg Gly Gly Gln Ile Met Thr Leu Lys Arg Asp Glu Thr 2655 2660 2665	8200
ATC CAG GAT GGC TGT GAC AGT CAC TTC TGC AAG GTC AAT GAA AGA GGA Ile Gln Asp Gly Cys Asp Ser His Phe Cys Lys Val Asn Glu Arg Gly 2670 2675 2680	8248
GAG TAC ATC TGG GAG AAG AGA GTC ACG GGT TGC CCA CCT TTC GAT GAA Glu Tyr Ile Trp Glu Lys Arg Val Thr Gly Cys Pro Pro Phe Asp Glu 2685 2690 2695	8296
CAC AAG TGT CTG GCT GAG GGA GGA AAA ATC ATG AAA ATT CCA GGC ACC His Lys Cys Leu Ala Glu Gly Lys Ile Met Lys Ile Pro Gly Thr 2700 2705 2710	8344
TGC TGT GAC ACA TGT GAG GAG CCA GAA TGC AAG GAT ATC ATT GCC AAG Cys Cys Asp Thr Cys Glu Glu Pro Glu Cys Lys Asp Ile Ile Ala Lys 2715 2720 2725 2730	8392
CTG CAG CGT GTC AAA GTG GGA GAC TGT AAG TCT GAA GAG GAA GTG GAC Leu Gln Arg Val Lys Val Gly Asp Cys Lys Ser Glu Glu Glu Val Asp 2735 2740 2745	8440
ATT CAT TAC TGT GAG GGT AAA TGT GCC AGC AAA GCC GTG TAC TCC ATC Ile His Tyr Cys Glu Gly Lys Cys Ala Ser Lys Ala Val Tyr Ser Ile 2750 2755 2760	8488
CAC ATG GAG GAT GTG CAG GAC CAG TGC TCC TGC TGC TCG CCC ACC CAG His Met Glu Asp Val Gln Asp Gln Cys Ser Cys Cys Ser Pro Thr Gln 2765 2770 2775	8536
ACG GAG CCC ATG CAG GTG GCC CTG CGC TGC ACC AAT GGC TCC CTC ATC Thr Glu Pro Met Gln Val Ala Leu Arg Cys Thr Asn Gly Ser Leu Ile 2780 2785 2790	8584
TAC CAT GAG ATC CTC AAT GCC ATC GAA TGC AGG TGT TCC CCC AGG AAG Tyr His Glu Ile Leu Asn Ala Ile Glu Cys Arg Cys Ser Pro Arg Lys 2795 2800 2805 2810	8632
TGC AGC AAG TGAGGCCACT GCCTGGATGC TACTGTCGCC TGCCTTACCC Cys Ser Lys	8681
GACCTCACTG GACTGGCCAG AGTGCTGCTC AGTCCTCCTC AGTCCTCCTC CTGCTCTGCT	8741
CTTGTGCTTC CTGATCCCAC AATAAAGGTC AATCTTCAC CTTGAAAAAAA AAAAAAAA	8801
A	8802

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2813 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Ser Pro Thr Arg Leu Val Arg Val Leu Leu Ala Leu Ala Leu Ile
1 5 10 15

Leu Pro Gly Lys Leu Cys Thr Lys Gly Thr Val Gly Arg Ser Ser Met
20 25 30

Ala Arg Cys Ser Leu Leu Gly Gly Asp Phe Ile Asn Thr Phe Asp Glu
35 40 45

Ser Met Tyr Ser Phe Ala Gly Asp Cys Ser Tyr Leu Leu Ala Gly Asp
50 55 60

Cys Gln Glu His Ser Ile Ser Leu Ile Gly Gly Phe Gln Asn Asp Lys
65 70 75 80

Arg Val Ser Leu Ser Val Tyr Leu Gly Glu Phe Phe Asp Ile His Leu
85 90 95

Phe Val Asn Gly Thr Met Leu Gln Gly Thr Gln Ser Ile Ser Met Pro
100 105 110

Tyr Ala Ser Asn Gly Leu Tyr Leu Glu Ala Glu Ala Gly Tyr Tyr Lys
115 120 125

Leu Ser Ser Glu Ala Tyr Gly Phe Val Ala Arg Ile Asp Gly Asn Gly
130 135 140

Asn Phe Gln Val Leu Leu Ser Asp Arg Tyr Phe Asn Lys Thr Cys Gly
145 150 155 160

Leu Cys Gly Asn Phe Asn Ile Phe Ala Glu Asp Asp Phe Lys Thr Gln
165 170 175

Glu Gly Thr Leu Thr Ser Asp Pro Tyr Asp Phe Ala Asn Ser Trp Ala
180 185 190

Leu Ser Ser Gly Glu Gln Arg Cys Lys Arg Val Ser Pro Pro Ser Ser
195 200 205

Pro Cys Asn Val Ser Ser Asp Glu Val Gln Gln Val Leu Trp Glu Gln
210 215 220

Cys Gln Leu Leu Lys Ser Ala Ser Val Phe Ala Arg Cys His Pro Leu
225 230 235 240

Val Asp Pro Glu Pro Phe Val Ala Leu Cys Glu Arg Thr Leu Cys Thr
245 250 255

Cys Val Gln Gly Met Glu Cys Pro Cys Ala Val Leu Leu Glu Tyr Ala
260 265 270

Arg Ala Cys Ala Gln Gln Gly Ile Val Leu Tyr Trp Thr Asp His
275 280 285

Ser Val Cys Arg Pro Ala Cys Pro Ala Gly Met Glu Tyr Lys Glu Cys
290 295 300

Val Ser Pro Cys Thr Arg Thr Cys Gln Ser Leu His Val Lys Glu Val
305 310 315 320

Cys Gln Glu Gln Cys Val Asp Gly Cys Ser Cys Pro Glu Gly Gln Leu
325 330 335

Leu Asp Glu Gly His Cys Val Gly Ser Ala Glu Cys Ser Cys Val His
340 345 350

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Ala Gly Gln Arg Tyr Pro Pro Gly Ala Ser Leu Leu Gln Asp Cys His
 355 360 365
 Thr Cys Ile Cys Arg Asn Ser Leu Trp Ile Cys Ser Asn Glu Glu Cys
 370 375 380
 Pro Gly Glu Cys Leu Val Thr Gly Gln Ser His Phe Lys Ser Phe Asp
 385 390 395 400
 Asn Arg Tyr Phe Thr Phe Ser Gly Val Cys His Tyr Leu Leu Ala Gln
 405 410 415
 Asp Cys Gln Asp His Thr Phe Ser Val Val Ile Glu Thr Val Gln Cys
 420 425 430
 Ala Asp Asp Leu Asp Ala Val Cys Thr Arg Ser Val Thr Val Arg Leu
 435 440 445
 Pro Gly His His Asn Ser Leu Val Lys Leu Lys Asn Gly Gly Val
 450 455 460
 Ser Met Asp Gly Gln Asp Ile Gln Ile Pro Leu Leu Gln Gly Asp Leu
 465 470 475 480
 Arg Ile Gln His Thr Val Met Ala Ser Val Arg Leu Ser Tyr Gly Glu
 485 490 495
 Asp Leu Gln Met Asp Ser Asp Val Arg Gly Arg Leu Leu Val Thr Leu
 500 505 510
 Tyr Pro Ala Tyr Ala Gly Lys Thr Cys Gly Arg Gly Asn Tyr Asn
 515 520 525
 Gly Asn Arg Gly Asp Asp Phe Val Thr Pro Ala Gly Leu Ala Glu Pro
 530 535 540
 Leu Val Glu Asp Phe Gly Asn Ala Trp Lys Leu Leu Gly Ala Cys Glu
 545 550 555 560
 Asn Leu Gln Lys Gln His Arg Asp Pro Cys Ser Leu Asn Pro Arg Gln
 565 570 575
 Ala Arg Phe Ala Glu Glu Ala Cys Ala Leu Leu Thr Ser Ser Lys Phe
 580 585 590
 Glu Pro Cys His Arg Ala Val Gly Pro Gln Pro Tyr Val Gln Asn Cys
 595 600 605
 Leu Tyr Asp Val Cys Ser Cys Ser Asp Gly Arg Asp Cys Leu Cys Ser
 610 615 620
 Ala Val Ala Asn Tyr Ala Ala Val Ala Arg Arg Gly Val His Ile
 625 630 635 640
 Ala Trp Arg Glu Pro Gly Phe Cys Ala Leu Ser Cys Pro Gln Gly Gln
 645 650 655
 Val Tyr Leu Gln Cys Gly Thr Pro Cys Asn Met Thr Cys Leu Ser Leu
 660 665 670
 Ser Tyr Pro Glu Glu Asp Cys Asn Glu Val Cys Leu Glu Ser Cys Phe
 675 680 685
 Ser Pro Pro Gly Leu Tyr Leu Asp Glu Arg Gly Asp Cys Val Pro Lys
 690 695 700

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Ala Gln Cys Pro Cys Tyr Tyr Asp Gly Glu Ile Phe Gln Pro Glu Asp
 705 710 715 720
 Ile Phe Ser Asp His His Thr Met Cys Tyr Cys Glu Asp Gly Phe Met
 725 730 735
 His Cys Thr Thr Ser Gly Gly Leu Gly Ser Leu Leu Pro Asn Pro Val
 740 745 750
 Leu Ser Ser Pro Arg Cys His Arg Ser Lys Arg Ser Leu Ser Cys Arg
 755 760 765
 Pro Pro Met Val Lys Leu Val Cys Pro Ala Asp Asn Pro Arg Ala Glu
 770 775 780
 Gly Leu Glu Cys Ala Lys Thr Cys Gln Asn Tyr Asp Leu Gln Cys Met
 785 790 795 800
 Ser Thr Gly Cys Val Ser Gly Cys Leu Cys Pro Gln Gly Met Val Arg
 805 810 815
 His Glu Asn Arg Cys Val Ala Leu Glu Arg Cys Pro Cys Phe His Gln
 820 825 830
 Gly Gln Glu Tyr Ala Pro Gly Glu Thr Val Lys Ile Asp Cys Asn Thr
 835 840 845
 Cys Val Cys Arg Asp Arg Lys Trp Thr Cys Thr Asp His Val Cys Asp
 850 855 860
 Ala Thr Cys Ser Ala Ile Gly Met Ala His Tyr Leu Thr Phe Asp Gly
 865 870 875 880
 Leu Lys Tyr Leu Phe Pro Gly Glu Cys Gln Tyr Val Leu Val Gln Asp
 885 890 895
 Tyr Cys Gly Ser Asn Pro Gly Thr Leu Arg Ile Leu Val Gly Asn Glu
 900 905 910
 Gly Cys Ser Tyr Pro Ser Val Lys Cys Lys Lys Arg Val Thr Ile Leu
 915 920 925
 Val Glu Gly Gly Glu Ile Glu Leu Phe Asp Gly Glu Val Asn Val Lys
 930 935 940
 Lys Pro Met Lys Asp Glu Thr His Phe Glu Val Val Glu Ser Gly Gln
 945 950 955 960
 Tyr Val Ile Leu Leu Leu Gly Lys Ala Leu Ser Val Val Trp Asp His
 965 970 975
 Arg Leu Ser Ile Ser Val Thr Leu Lys Arg Thr Tyr Gln Glu Gln Val
 980 985 990
 Cys Gly Leu Cys Gly Asn Phe Asp Gly Ile Gln Asn Asn Asp Phe Thr
 995 1000 1005
 Ser Ser Ser Leu Gln Ile Glu Glu Asp Pro Val Asp Phe Gly Asn Ser
 1010 1015 1020
 Trp Lys Val Asn Pro Gln Cys Ala Asp Thr Lys Lys Val Pro Leu Asp
 1025 1030 1035 1040
 Ser Ser Pro Ala Val Cys His Asn Asn Ile Met Lys Gln Thr Met Val
 1045 1050 1055

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Asp Ser Ser Cys Arg Ile Leu Thr Ser Asp Ile Phe Gln Asp Cys Asn
 1060 1065 1070
 Arg Leu Val Asp Pro Glu Pro Phe Leu Asp Ile Cys Ile Tyr Asp Thr
 1075 1080 1085
 Cys Ser Cys Glu Ser Ile Gly Asp Cys Thr Cys Phe Cys Asp Thr Ile
 1090 1095 1100
 Ala Ala Tyr Ala His Val Cys Ala Gln His Gly Lys Val Val Ala Trp
 1105 1110 1115 1120
 Arg Thr Ala Thr Phe Cys Pro Gln Asn Cys Glu Glu Arg Asn Leu His
 1125 1130 1135
 Glu Asn Gly Tyr Glu Cys Glu Trp Arg Tyr Asn Ser Cys Ala Pro Ala
 1140 1145 1150
 Cys Pro Ile Thr Cys Gln His Pro Glu Pro Leu Ala Cys Pro Val Gln
 1155 1160 1165
 Cys Val Glu Gly Cys His Ala His Cys Pro Pro Gly Lys Ile Leu Asp
 1170 1175 1180
 Glu Leu Leu Gln Thr Cys Ile Asp Pro Glu Asp Cys Pro Val Cys Glu
 1185 1190 1195 1200
 Val Ala Gly Arg Arg Leu Ala Pro Gly Lys Lys Ile Ile Leu Asn Pro
 1205 1210 1215
 Ser Asp Pro Glu His Cys Gln Ile Cys Asn Cys Asp Gly Val Asn Phe
 1220 1225 1230
 Thr Cys Lys Ala Cys Arg Glu Pro Gly Ser Val Val Pro Pro Thr
 1235 1240 1245
 Asp Gly Pro Ile Gly Ser Thr Thr Ser Tyr Val Glu Asp Thr Ser Glu
 1250 1255 1260
 Pro Pro Leu His Asp Phe His Cys Ser Arg Leu Leu Asp Leu Val Phe
 1265 1270 1275 1280
 Leu Leu Asp Gly Ser Ser Lys Leu Ser Glu Asp Glu Phe Glu Val Leu
 1285 1290 1295
 Lys Val Phe Val Val Gly Met Met Glu His Leu His Ile Ser Gln Lys
 1300 1305 1310
 Arg Ile Arg Val Ala Val Val Glu Tyr His Asp Gly Ser His Ala Tyr
 1315 1320 1325
 Ile Glu Leu Lys Asp Arg Lys Arg Pro Ser Glu Leu Arg Arg Ile Thr
 1330 1335 1340
 Ser Gln Val Lys Tyr Ala Gly Ser Glu Val Ala Ser Thr Ser Glu Val
 1345 1350 1355 1360
 Leu Lys Tyr Thr Leu Phe Gln Ile Phe Gly Lys Ile Asp Arg Pro Glu
 1365 1370 1375
 Ala Ser Arg Ile Ala Leu Leu Met Ala Ser Gln Glu Pro Ser Arg
 1380 1385 1390
 Leu Ala Arg Asn Leu Val Arg Tyr Val Gln Gly Leu Lys Lys Lys
 1395 1400 1405

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Val Ile Val Ile Pro Val Gly Ile Gly Pro His Ala Ser Leu Lys Gln
1410 1415 1420

Ile His Leu Ile Glu Lys Gln Ala Pro Glu Asn Lys Ala Phe Val Phe
1425 1430 1435 1440

Ser Gly Val Asp Glu Leu Glu Gln Arg Arg Asp Glu Ile Ile Asn Tyr
1445 1450 1455

Leu Cys Asp Leu Ala Pro Glu Ala Pro Ala Pro Thr Gln His Pro Pro
1460 1465 1470

Met Ala Gln Val Thr Val Gly Ser Glu Leu Leu Gly Val Ser Ser Pro
1475 1480 1485

Gly Pro Lys Arg Asn Ser Met Val Leu Asp Val Val Phe Val Leu Glu
1490 1495 1500

Gly Ser Asp Lys Ile Gly Glu Ala Asn Phe Asn Lys Ser Arg Glu Phe
1505 1510 1515 1520

Met Glu Glu Val Ile Gln Arg Met Asp Val Gly Gln Asp Arg Ile His
1525 1530 1535

Val Thr Val Leu Gln Tyr Ser Tyr Met Val Thr Val Glu Tyr Thr Phe
1540 1545 1550

Ser Glu Ala Gln Ser Lys Gly Glu Val Leu Gln Gln Val Arg Asp Ile
1555 1560 1565

Arg Tyr Arg Gly Gly Asn Arg Thr Asn Thr Gly Leu Ala Leu Gln Tyr
1570 1575 1580

Leu Ser Glu His Ser Phe Ser Val Ser Gln Gly Asp Arg Glu Gln Val
1585 1590 1595 1600

Pro Asn Leu Val Tyr Met Val Thr Gly Asn Pro Ala Ser Asp Glu Ile
1605 1610 1615

Lys Arg Met Pro Gly Asp Ile Gln Val Val Pro Ile Gly Val Gly Pro
1620 1625 1630

His Ala Asn Val Gln Glu Leu Glu Lys Ile Gly Trp Pro Asn Ala Pro
1635 1640 1645

Ile Leu Ile His Asp Phe Glu Met Leu Pro Arg Glu Ala Pro Asp Leu
1650 1655 1660

Val Leu Gln Arg Cys Cys Ser Gly Glu Gly Leu Gln Ile Pro Thr Leu
1665 1670 1675 1680

Ser Pro Thr Pro Asp Cys Ser Gln Pro Leu Asp Val Val Leu Leu
1685 1690 1695

Asp Gly Ser Ser Ser Ile Pro Ala Ser Tyr Phe Asp Glu Met Lys Ser
1700 1705 1710

Phe Thr Lys Ala Phe Ile Ser Arg Ala Asn Ile Gly Pro Arg Leu Thr
1715 1720 1725

Gln Val Ser Val Leu Gln Tyr Gly Ser Ile Thr Thr Ile Asp Val Pro
1730 1735 1740

Trp Asn Val Ala Tyr Glu Lys Val His Leu Leu Ser Leu Val Asp Leu
1745 1750 1755 1760

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Met Gln Gln Glu Gly Gly Pro Ser Glu Ile Gly Asp Ala Leu Ser Phe
 1765 1770 1775
 Ala Val Arg Tyr Val Thr Ser Glu Val His Gly Ala Arg Pro Gly Ala
 1780 1785 1790
 Ser Lys Ala Val Val Ile Leu Val Thr Asp Val Ser Val Asp Ser Val
 1795 1800 1805
 Asp Ala Ala Ala Glu Ala Ala Arg Ser Asn Arg Val Thr Val Phe Pro
 1810 1815 1820
 Ile Gly Ile Gly Asp Arg Tyr Ser Glu Ala Gln Leu Ser Ser Leu Ala
 1825 1830 1835 1840
 Gly Pro Lys Ala Gly Ser Asn Met Val Arg Leu Gln Arg Ile Glu Asp
 1845 1850 1855
 Leu Pro Thr Val Ala Thr Leu Gly Asn Ser Phe Phe His Lys Leu Cys
 1860 1865 1870
 Ser Gly Phe Asp Arg Val Cys Val Asp Glu Asp Gly Asn Glu Lys Arg
 1875 1880 1885
 Pro Gly Asp Val Trp Thr Leu Pro Asp Gln Cys His Thr Val Thr Cys
 1890 1895 1900
 Leu Pro Asp Gly Gln Thr Leu Leu Lys Ser His Arg Val Asn Cys Asp
 1905 1910 1915 1920
 Arg Gly Pro Arg Pro Ser Cys Pro Asn Gly Gln Pro Pro Leu Arg Val
 1925 1930 1935
 Glu Glu Thr Cys Gly Cys Arg Trp Thr Cys Pro Cys Val Cys Met Gly
 1940 1945 1950
 Ser Ser Thr Arg His Ile Val Thr Phe Asp Gly Gln Asn Phe Lys Leu
 1955 1960 1965
 Thr Gly Ser Cys Ser Tyr Val Leu Phe Gln Asn Lys Glu Gln Asp Leu
 1970 1975 1980
 Glu Val Ile Leu Gln Asn Gly Ala Cys Ser Pro Gly Ala Lys Glu Thr
 1985 1990 1995 2000
 Cys Met Lys Ser Ile Glu Val Lys His Asp Gly Leu Ser Val Glu Leu
 2005 2010 2015
 His Ser Asp Met Gln Met Thr Val Asn Gly Arg Leu Val Ser Ile Pro
 2020 2025 2030
 Tyr Val Gly Gly Asp Met Glu Val Asn Val Tyr Gly Thr Ile Met Tyr
 2035 2040 2045
 Glu Val Arg Phe Asn His Leu Gly His Ile Phe Thr Phe Thr Pro Gln
 2050 2055 2060
 Asn Asn Glu Phe Gln Leu Gln Leu Ser Pro Arg Thr Phe Ala Ser Lys
 2065 2070 2075 2080
 Thr Tyr Gly Leu Cys Gly Ile Cys Asp Glu Asn Gly Ala Asn Asp Phe
 2085 2090 2095
 Ile Leu Arg Asp Gly Thr Val Thr Asp Trp Lys Ala Leu Ile Gln
 2100 2105 2110

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Glu Trp Thr Val Gln Gln Leu Gly Lys Thr Ser Gln Pro Val His Glu
2115 2120 2125

Glu Gln Cys Pro Val Ser Glu Phe Phe His Cys Gln Val Leu Leu Ser
2130 2135 2140

Glu Leu Phe Ala Glu Cys His Lys Val Leu Ala Pro Ala Thr Phe Tyr
2145 2150 2155 2160

Ala Met Cys Gln Pro Asp Ser Cys His Pro Lys Lys Val Cys Glu Ala
2165 2170 2175

Ile Ala Leu Tyr Ala His Leu Cys Arg Thr Lys Gly Val Cys Val Asp
2180 2185 2190

Trp Arg Arg Ala Asn Phe Cys Ala Met Ser Cys Pro Pro Ser Leu Val
2195 2200 2205

Tyr Asn His Cys Glu His Gly Cys Pro Arg Leu Cys Glu Gly Asn Thr
2210 2215 2220

Ser Ser Cys Gly Asp Gln Pro Ser Glu Gly Cys Phe Cys Pro Pro Asn
2225 2230 2235 2240

Gln Val Met Leu Glu Gly Ser Cys Val Pro Glu Glu Ala Cys Thr Gln
2245 2250 2255

Cys Ile Ser Glu Asp Gly Val Arg His Gln Phe Leu Glu Thr Trp Val
2260 2265 2270

Pro Ala His Gln Pro Cys Gln Ile Cys Thr Cys Leu Ser Gly Arg Lys
2275 2280 2285

Val Asn Cys Thr Leu Gln Pro Cys Pro Thr Ala Lys Ala Pro Thr Cys
2290 2295 2300

Gly Pro Cys Glu Val Ala Arg Leu Arg Gln Asn Ala Val Gln Cys Cys
2305 2310 2315 2320

Pro Glu Tyr Glu Cys Val Cys Asp Leu Val Ser Cys Asp Leu Pro Pro
2325 2330 2335

Val Pro Pro Cys Glu Asp Gly Leu Gln Met Thr Leu Thr Asn Pro Gly
2340 2345 2350

Glu Cys Arg Pro Asn Phe Thr Cys Ala Cys Arg Lys Asp Glu Cys Arg
2355 2360 2365

Arg Glu Ser Pro Pro Ser Cys Pro Pro His Arg Thr Pro Ala Leu Arg
2370 2375 2380

Lys Thr Gln Cys Cys Asp Glu Tyr Glu Cys Ala Cys Asn Cys Val Asn
2385 2390 2395 2400

Ser Thr Val Ser Cys Pro Leu Gly Tyr Leu Ala Ser Ala Val Thr Asn
2405 2410 2415

Asp Cys Gly Cys Thr Thr Cys Phe Pro Asp Lys Val Cys Val
2420 2425 2430

His Arg Gly Thr Ile Tyr Pro Val Gly Gln Phe Trp Glu Glu Ala Cys
2435 2440 2445

Asp Val Cys Thr Cys Thr Asp Leu Glu Asp Ser Val Met Gly Leu Arg
2450 2455 2460

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Val Ala Gln Cys Ser Gln Lys Pro Cys Glu Asp Asn Cys Leu Ser Gly
 2465 2470 2475 2480
 Phe Thr Tyr Val Leu His Glu Gly Glu Cys Cys Gly Arg Cys Leu Pro
 2485 2490 2495
 Ser Ala Cys Glu Val Val Thr Gly Ser Pro Arg Gly Asp Ala Gln Ser
 2500 2505 2510
 His Trp Lys Asn Val Gly Ser His Trp Ala Ser Pro Asp Asn Pro Cys
 2515 2520 2525
 Leu Ile Asn Glu Cys Val Arg Val Lys Glu Glu Val Phe Val Gln Gln
 2530 2535 2540
 Arg Asn Val Ser Cys Pro Gln Leu Asn Val Pro Thr Cys Pro Thr Gly
 2545 2550 2555 2560
 Phe Gln Leu Ser Cys Lys Thr Ser Glu Cys Cys Pro Thr Cys His Cys
 2565 2570 2575
 Glu Pro Leu Glu Ala Cys Leu Leu Asn Gly Thr Ile Ile Gly Pro Gly
 2580 2585 2590
 Lys Ser Leu Met Ile Asp Val Cys Thr Thr Cys Arg Cys Thr Val Pro
 2595 2600 2605
 Val Gly Val Ile Ser Gly Phe Lys Leu Glu Gly Arg Lys Thr Thr Cys
 2610 2615 2620
 Glu Ala Cys Pro Leu Gly Tyr Lys Glu Glu Lys Asn Gln Gly Glu Cys
 2625 2630 2635 2640
 Cys Gly Arg Cys Leu Pro Ile Ala Cys Thr Ile Gln Leu Arg Gly Gly
 2645 2650 2655
 Gln Ile Met Thr Leu Lys Arg Asp Glu Thr Ile Gln Asp Gly Cys Asp
 2660 2665 2670
 Ser His Phe Cys Lys Val Asn Glu Arg Gly Glu Tyr Ile Trp Glu Lys
 2675 2680 2685
 Arg Val Thr Gly Cys Pro Pro Phe Asp Glu His Lys Cys Leu Ala Glu
 2690 2695 2700
 Gly Gly Lys Ile Met Lys Ile Pro Gly Thr Cys Cys Asp Thr Cys Glu
 2705 2710 2715 2720
 Glu Pro Glu Cys Lys Asp Ile Ile Ala Lys Leu Gln Arg Val Lys Val
 2725 2730 2735
 Gly Asp Cys Lys Ser Glu Glu Val Asp Ile His Tyr Cys Glu Gly
 2740 2745 2750
 Lys Cys Ala Ser Lys Ala Val Tyr Ser Ile His Met Glu Asp Val Gln
 2755 2760 2765
 Asp Gln Cys Ser Cys Cys Ser Pro Thr Gln Thr Glu Pro Met Gln Val
 2770 2775 2780
 Ala Leu Arg Cys Thr Asn Gly Ser Leu Ile Tyr His Glu Ile Leu Asn
 2785 2790 2795 2800
 Ala Ile Glu Cys Arg Cys Ser Pro Arg Lys Cys Ser Lys
 2805 2810

WE CLAIM:

1. An isolated nucleic acid comprising a nucleotide sequence encoding canine von Willebrand Factor polypeptide.
2. The isolated nucleic acid of Claim 1, wherein the nucleotide sequence is capable of hybridizing under high stringency conditions to SEQ ID NO. 1.
3. The isolated nucleic acid of Claim 1, wherein the nucleotide sequence encodes the Scottish terrier von Willebrand Factor polypeptide.
4. The isolated nucleic acid of Claim 2, wherein the nucleotide sequence encodes the Scottish terrier von Willebrand Factor polypeptide.
- 10 5. A vector comprising the nucleic acid of Claim 1.
6. A vector comprising the nucleic acid of Claim 2.
7. A cell comprising the vector of Claim 5.
8. A cell comprising the vector of Claim 6.
9. An isolated nucleic acid comprising a nucleotide sequence encoding 15 defective canine von Willebrand Factor polypeptide.
10. The isolated nucleic acid of Claim 9, wherein the nucleotide sequence is capable of hybridizing under high stringency conditions to the complement of SEQ ID NO. 1 having a base deletion at codon 88.
11. A vector comprising the nucleic acid of Claim 9.
- 20 12. A vector comprising the nucleic acid of Claim 10.
13. A cell comprising the vector of Claim 11.
14. A cell comprising the vector of Claim 12.

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15. An isolated oligonucleotide sequence consisting of contiguous nucleic acids of the nucleotide sequence of SEQ ID NO. 1 and capable of specifically hybridizing with the canine von Willebrand Factor gene.

16. An isolated oligonucleotide sequence consisting of contiguous nucleic acids of the nucleotide sequence that is complementary to the sequence of SEQ ID NO. 1 and capable of specifically hybridizing with the canine von Willebrand Factor gene.

17. A method of detecting a canine von Willebrand Factor gene in a sample comprising the steps of:

10 a) contacting the sample with a oligonucleotide comprising contiguous nucleic acids of the nucleotide sequence of SEQ ID NO. 1 and capable of specifically hybridizing with the canine von Willebrand Factor gene, under conditions favorable for hybridization of the oligonucleotide to any complementary sequences of nucleic acid in the sample; and

15 b) detecting hybridization, thereby detecting a canine von Willebrand Factor gene.

18. The method of Claim 17, further comprising the step of:

20 c) quantifying hybridization of the oligonucleotide to complementary sequence.

19. The method of Claim 17, wherein in SEQ ID NO. 1 there is a base deletion at codon 88.

20. An assay kit for screening for a canine von Willebrand Factor gene comprising:

25 a) an oligonucleotide comprising contiguous nucleic acids of the nucleotide sequence of SEQ ID NO. 1 and capable of hybridizing with the canine von Willebrand Factor gene;

b) reagents for hybridization of the oligonucleotide to a complementary nucleic acid sequence; and

30 c) container means for a)-b).

21. A method of detecting a canine von Willebrand Factor gene in a sample comprising the steps of:

- 5 a) contacting the sample with an oligonucleotide comprising contiguous nucleic acids of the nucleotide sequence that is complementary to the sequence of SEQ ID NO. 1 and capable of specifically hybridizing to the complementary nucleotide sequence, under conditions favorable for hybridization of the oligonucleotide to any complementary sequences of nucleic acid in the sample; and
- 10 b) detecting hybridization, thereby detecting a canine von Willebrand Factor gene.

22. The method of Claim 21, further comprising the step of:

- c) quantifying hybridization of the oligonucleotide to complementary sequences.

15 23. The method of Claim 21, wherein in SEQ ID NO. 1 there is a base deletion at codon 88.

24. An assay kit for screening for a canine von Willebrand Factor gene comprising:

- 20 a) an oligonucleotide comprising contiguous acids from the nucleotide sequence that is complementary to the sequence of SEQ ID NO. 1 and capable of specifically hybridizing to the complementary nucleotide sequence;
- b) reagents for hybridization of the oligonucleotide to a complementary nucleic acid sequence; and
- 25 c) container means for a)-b).

25. The assay kit of Claim 24, wherein in SEQ ID NO. 1 there is a base deletion at codon 88.

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26. A method for detecting a mutated canine von Willebrand Factor gene in a canine DNA sample comprising the steps of:

- 5 a) amplifying the DNA sample by polymerase chain reaction to produce polymerase chain reaction products, wherein the polymerase chain reaction uses primers that produce a restriction site in a mutant allele but not in a normal allele;
- b) digesting the polymerase chain reaction products with a restriction enzyme specific to the restriction site of the restriction site primer to produce DNA fragments; and
- 10 c) detecting the DNA fragments, thereby detecting a mutated canine von Willebrand Factor gene.

27. The method of Claim 26, wherein the primers are those of Figure 4.

28. The method of Claim 26, wherein the DNA fragments are detected by gel electrophoresis.

15 29. The method of Claim 27, wherein the restriction enzyme is *Bsi*EI.

30. The method of Claim 27, wherein the restriction enzyme is *Sau*96 I.

31. An oligonucleotide probe capable of detecting a mutation associated with canine von Willebrand's disease, wherein the mutation is a base deletion at codon 88 of the canine von Willebrand Factor gene.

FIGURE 1A

1 CATTAAANAGG TCCTGGCTGG GAGCTTTTT TTGGGACCAAG CACTCCATGT TCAAGGGCAA
 61 ACAGGGGCCA ATTAGGATCA ATCTTTTTC TTTCTTTTT TAAAAAAA AATTCTTCCC
 121 ACTTTGCACA CGGACAGTAG TACATACCAAG TAGCTCTCTG CGAGGACGGT GATCACTAAT
 181 CATTCTCCT GCTTCGTGGC AGATGAGTCC TACCAAGACTT GTGAGGGTGC TGCTGGCTCT
 241 GCCCTCATC TTGCCAGGG AACTTTGTAC AAAAGGGACT GTTGAAGGTT CATCGATGGC
 301 CCGATGTAGC CTCTCGGAG GTGACTTCAT CAACACCTT GATGAGAGCA TGTACAGCTT
 361 TGCAGGGAGAT TGCAGTTACC TCTGGCTGG GGACTGCCAG GAAACACTCCA TCTCACTTAT
 421 CGGGGGTTTC CAAAATGACA AAAGAGTGAAG CCTCTCCGTG TATCTCGGAG AATTTTTCGA
 481 CATTCAATTG TTTGTCATG GTACCATGCT GCAGGGGACCA CAAAGCATCT CCATGCCCTA
 541 CGCCCTCAAAT GGGCTGTATC TAGAGGCCGA GGCTGGCTAC TACAAGCTGT CCAGTGAGGC
 601 CTACGGCTTT GTGGCCAGAA TTGATGGCAA TGGCAACTTT CAAGTCTCTG TGTCAGACAG
 661 ATACTTCAAC AAGACCTGTG GGCTGTGTGG CAACCTTAAT ATCTTTGCTG AGGATGACTT
 721 CAAGACTCAA GAAGGGACGT TGACTTCGGA CCCCTATGAC TTTGCCAACT CCTGGGCCCT
 781 GAGCAGTGGG GAACAACGGT GCAAACGGGT GTCCCCCTCCC AGCAGCCCCAT GCAATGTCTC
 841 CTCTGATGAA GTGCAGCAGG TCTCTGGGA GCAGTGCCAG CTCCTGAAGA GTGCCCTCGGT
 901 GTTTGCCCGC TGCCACCCGC TGGTGGACCC TGAGCCTTTT GTGCCCTCTG TGAAAGGAC
 961 TCTGTGCACC TGTGTCCAGG GGATGGAGTG CCCCTGTGCG GTCCCTCCTGG AGTACGCCCG
 1021 GCCCTGTGCC CAGCAGGGGA TTGTCCTTGTG CCGCTGGAC GACCACAGCG TCTGCCGACC
 1081 AGCATGCCCT GCTGGCATGG AGTACAAGGA GTGCGTGTCC CTTGCACCA GAACTTGCCA
 1141 GAGCCTTCAT GTCAAAGAAG TGTGTCAAGG GCAATGTGTA GATGGCTGCA GCTGCCCGA
 1201 GGGCCAGCTC CTGGATGAAG GCAACTGCGT GGGAAAGTGCT GAGTGTTCCT GTGTGCATGC
 1261 TGGGCAACGG TACCCCTCCGG GCGCCTCCCT CTTACAGGAC TGCCACACCT GCATTTGCCG
 1321 AAATAGCCCTG TGGATCTGCA GCAATGAAGA ATGCCACAGGC GAGTGTCTGG TCACAGGACA
 1381 GTCCCACCTC AAGAGCTTCG ACAACAGGTG CTTCACCTTC AGTGGGGTCT GCCACTACCT
 1441 GCTGGCCCAAG GACTGCCAGG ACCACACATT CTCTGTGTGTC ATAGAGACTG TCCAGTGTGC
 1501 CGATGACCTG GATGCTGTCT GCACCCGCTC GGTACCGTC CGCCTGCCTG GACATCACAA
 1561 CAGCCTTGTG AAGCTGAAGA ATGGGGGAGG AGTCTCCATG GATGGCCAGG ATATCCAGAT
 1621 TCTCTCCTG CAAGGTGACC TCCGCATCCA GCACACCGTG ATGGCCTCCG TGCGCCTCAG
 1681 CTACGGGGAG GACCTGCAGA TGGATTGCGA CGTCCCCGGC AGGCTACTGG TGACGCTGTA
 1741 CCCCCTCAG GCGGGGAAGA CGTGCAGGGCG TGGCGGGAAAC TACAACGGCA ACCGGGGGGA
 1801 CGACTTCGTG ACGCCCGCAG GCCTGGCGGA GCCCCCTGGTG GAGGACTTCG GGAACGCCCTG
 1861 GAAGCTGCTC GGGGCTGCG AGAACCTGCA GAAGCAGCAC CGCGATCCCT GCAGCCTCAA
 1921 CCCGCAGG GCAAGGTGTG CGGAGGAGGC GTGCGCGCTG CTGACGTTCT CGAAGTTGCA
 1981 GCCCCGCCAC CGAGCGGTGG GTCCCTCAGCC CTACGTGCAG AACTGCCCTCT ACGACGTCTG
 2041 CTCTGCTCC GACGGCAGAG ACTGTCTTG CAGCGCCGTG GCCAACCTACG CCGCAGCCGT
 2101 GGGCCGGAGG GGCCTGCACA TCGCGTGGCG GGAGCCGGGC TTCTGTGCGC TGAGCTGCC
 2161 CCAGGGCCAG GTGTACCTGC AGTGTGGGAC CCCCTGCAAC ATGACCTGTC TCTCCCTCTC
 2221 TTACCCGGAG GAGGACTGCA ATGAGGTCTG CTTGGAAAGC TGCTCTCTCC CCCCAGGGCT
 2281 GTACCTGGAG GAGAGGGAG ATTGTGTGCC CAAGGCTCAG TGTCCTGTGTT ACTATGATGG
 2341 TGAGATCTT CAGCCCAGG ACATCTTCTC AGACCATCAC ACCATGTGCT ACTGTGAGGA
 2401 TGGCTTCATG CACTGTACCA CAAGTGGAGG CCTGGGAAGC CTGCTGCCCA ACCCGGTGCT
 2461 CAGCAGCCCC CGGTGTCAACC GCAGCAAAG GAGCCTGTCC TGTCGGCCCC CCATGGCTAA
 2521 GTTGGTGTGT CCCCTGTATA ACCCGAGGGC TGAAGGACTG GAGTGTGCCA AAACCTGCCA
 2581 GAACTATGAC CTGCACTGCA TGAGCACAGG CTGTGTCTCC GGCTGCCTCT GCCCGCAGGG
 2641 CATGGTCAGG CATGAAAACA GGTGTGTGGC GCTGGAAAGA TGTCCTGTGCT TCCACCAAGG
 2701 CCAAGAGTAC GCCCCCAGGAG AAACCGTGAAG AATTGACTGC AACACTTGAG TCTGTGGGA
 2761 CGCGAAGTGG ACCTGCACAG ACCATGTGTG TGATGCCACT TGCTCTGCCA TCGGCATGGC
 2821 GCACTACCTC ACCTTCGACG GACTCAAGTA CCTGTCCCT GGGGAGTGCC AGTATGTTCT
 2881 GGTGCAGGAT TACTGCAGGCA GTAAACCTGG GACCTTACGG ATCCCTGGTGG GGAACGGAGGG
 2941 GTGCAGCTAC CCCTCAGTGA AATGCAAGAA GCGGGTCACC ATCCCTGGTGG AAGGAGGAGA
 3001 GATTGAACGTG TTTGATGGGG AGGTGAATGT GAAGAAACCC ATGAAGGATG AGACTCACTT
 3061 TGAGGTGGTA GAGTGTGGTC AGTACGTCA TCTGTGCTG GGCAAGGCAC TCTCTGTGGT
 3121 CTGGGACAC CGCCTGAGCA TCTCTGTGAC CCTGAAGCGG ACATACCAGG AGCAGGTGTG

FIGURE 1B

3181 TGGCCTGTGT GGGAAATTTG ATGGCATCCA GAACAATGAT TTCAACCAGCA GCAGCCTCCA
 3241 AATAGAAGAA GACCCCTGTGG ACTTTGGGAA TTCTGGAAA GTGAACCCGC AGTGTGCCGA
 3301 CACCAAGAAA GTACCACTGG ACTCATCCCC TGCCTCTGC CACAACAACA TCATGAAGCA
 3361 GACGATGGTG GATTCCTCCCT GCAGGATCCT CACCAAGTGAT ATTTTCCAGG ACTGCAACAG
 3421 GCTGGTGGAC CCTGAGCCAT TCCTGGACAT TTGCATCTAC GACACTTGCT CCTGTGAGTC
 3481 CATTGGGGAC TGCACCTGCT TCTGTGACAC CATTGCTGCT TACGCCACG TCTGTGCCA
 3541 GCATGGCAAG GTGGTAGCCT GGAGGACAGC CACATTCTGT CCCCAGAATT GCGAGGAGCG
 3601 GAATCTCCAC GAGAATGGGT ATGAGTGTGA GTGGCGCTAT AACAGCTGTG CCCCTGCCGT
 3661 TCCCCTACAG TGCCAGCACC CCGAGCCACT GGCACTGCCCT GTACAGTGTG TTGAAGGTTG
 3721 CCATGCGCAC TGCCCTCCAG GGAAAATCCT GGATGAGCTT TTGCAGACT GCATGACCC
 3781 TGAAGACTGT CCTGTGTGTG AGGTGGTGG TCGTCGCTTG GCCCCAGGAA AGAAAATCAT
 3841 CTTGAACCCC AGTGACCCCTG AGCACTGCCA AATTTGTAAT TGTGATGGTG TCAACTTCAC
 3901 CTGTAAGGCC TGCAGAGAAC CCGGAAGTGT TGTGGTGCCT CCCACAGATG GCCCCATTGG
 3961 CTCTTACCAAC TCGTATGTGG AGGACACGTC GGAGCCGCCCT CTCCATGACT TCCACTGCAG
 4021 CAGGCTTCTG GACCTGGTTT TCCTGCTGGA TGGCTCCTCC AAGCTGTCTG AGGACGAGTT
 4081 TGAAGTGCTG AAGGTCTTGTG TGGTGGTAT GATGGAGCAT CTGCACATCT CCCAGAAGCG
 4141 GATCCCGTG GCTGTGGTGG AGTACCAACGA CGGCTCCAC GCCTACATCG AGCTCAAGGA
 4201 CCGGAAGCGA CCCTCAGAGC TGCAGCCAT CACCAAGCCAG GTGAAGTACG CGGGCAGCGA
 4261 GGTGGCCTCC ACCAGTGAGG TCTTAAAGTA CACGCTGTTC CAGATCTTGTG GCAAGATCGA
 4321 CGCCCCGAA GCGTCTCGCA TTGCCCCGTCT CCGTATGCC AGCCAGGAGC CCTCAAGGCT
 4381 GGCCCCGAA TTGGTCCGCT ATGTGCAAGGG CCTGAAGAAG AAGAAAGTCA TTGTCACTCCC
 4441 TGTGGGCATC GGGCCCCACCG CCAGCTTAA GCAGATCCAC CTCATAGAGA AGCAGGGCCC
 4501 TGAGAACAAG GCCTTTGTGT TCAGTGGTGT GGATGAGTTG GAGCAGCGA GGGATGAGAT
 4561 TATCAACTAC CTCTGTGACC TTGCCCCGA AGCACCTGCC CCTACTCAGC ACCCCCCAAT
 4621 GGCCCCAGGTC ACGGTGGTT CGGAGCTGTT GGGGGTTCA TCTCCAGGAC CCAAAAGGAA
 4681 CTCCATGGTC CTGGATGTGG TGTGGTCTCT GGAAGGGTCA GACAAAATTG GTGAGGCCA
 4741 CTTAACAAA AGCAGGGAGT TCATGGAGGA GGTGATTCA CGGATGGACG TGGGCCAGGA
 4801 CAGGATCCAC GTCACAGTGC TGCACTACTC GTACATGGTG ACCGTGGAGT ACACCTTCAG
 4861 CGAGGCGCAG TCCAAGGGCG AGGTCTTACA GCAGGTGCAG GATATCCGAT ACCGGGGTGG
 4921 CAACAGGACC AACACTGGAC TGGCCCTGCA ATACCTGTCC GAACACAGCT TCTCCGTCAG
 4981 CCAGGGGGAC CGGGAGCGAG TACCTAACCT GGTCTACATG GTCACAGGAA ACCCCCCTTC
 5041 TGATGAGATC AAGCGGATGC CTGGAGACAT CCAGGTGGTG CCCATCGGGG TGGGTCACAA
 5101 TGCCCAATGTG CAGGAGCTGG AGAAGATTGG CTGGCCCAAT GCCCCCATCC TCATCCATGA
 5161 CTTTGAGATG CTCCCTCGAG AGGCTCTGA TCTGGTGTCA CAGAGGTGCT GCTCTGGAGA
 5221 GGGGCTGCAG ATCCCCACCC TCTCCCCACCC CCCAGATTGC AGCCAGCCCC TGGATGTGGT
 5281 CCTCCCTCTG GATGGCTCTT CCAGCATTC AGCTTCTTAC TTTGATGAA TGAAGAGCTT
 5341 CACCAAGGCT TTTATTCAG GAGCTAATAT AGGGCCCCGG CTCACACTCAG TGTGGTGTCT
 5401 GCAATATGGA AGCATCACCA CTATCGATGT GCCTTGAAT GTAGCTATG AGAAAGTCCA
 5461 TTTACTGAGC CTTGTGGACC TCATGCAGCA GGAGGGAGGC CCCAGCGAAA TTGGGGATGC
 5521 TTTGAGCTT GCGTGCAGAT ATGTCACCTC AGAAGTCCAT GGTGCCAGGC CCGGAGCCTC
 5581 GAAAGCGGTG GTTATCCTAG TCACAGATGT CTCCGTGGAT TCAGTGGATG CTGCAGCGA
 5641 GGCCGCCAGA TCCAACCGAG TGACAGTGTG TCCCATTGGA ATCGGGGATC GGTACAGTGA
 5701 GGCCCAAGCTG AGCAGCTTGG CAGGCCAAA GGCTGGCTCC AATATGGTAA GGCTCCAGCG
 5761 AATTGAAGAC CTCCCCACCG TGGCCACCCCT GGGAAATTCC TTCTTCCACA AGCTGTGCTC
 5821 TGGGTTTGAT AGAGTTTGCG TGGATGAGGA TGGGAATGAG AAGAGGCCCG GGGATGTCTG
 5881 GACCTTGCA GACCACTGCCC ACACAGTGC TTGCTGCTCA GATGGCCAGA CCTTGCTGAA
 5941 GAGTCATCGG GTCAACTGTG ACCGGGGGCC AAGGCCCTCG TGCCCCAATG GCCAGCCCCC
 6001 TCTCAGGGTA GAGGAGACCT GTGGCTGCCG CTGGACCTGT CCCCTGTGTG GCATGGGCAG
 6061 CTCTACCCGG CACATCGTGA CCTTTGATGG GCAGAATTTC AAGCTGACTG GCAGCTGTTC
 6121 GTATGTCCTA TTTCAAAACA AGGAGCAGGA CCTGGAGGTG ATTCCTCAGA ATGGTGCCTG
 6181 CAGCCCTGGG GCGAAGGAGA CCTGCATGAA ATCCATTGAG GTGAAGCATG ACGGCCTCTC
 6241 AGTTGAGCTC CACAGTGACA TGCAGATGAC AGTGAATGGG AGACTAGTCT CCATCCCATA
 6301 TGTGGGTGGA GACATGGAAG TCAATGTTTA TGGGACCATC ATGTATGAGG TCAGATTCAA
 6361 CCATCTTGGC CACATCTTCA CATTCAACCC CCAAAACAAT GAGTTCCAGC TGCACTCAG

FIGURE 1C

6421 CCCCCAGGACC TTTGCTTCGA AGACATATGG TCTCTGTGGG ATCTGTGATG AGAACGGAGC
 6481 CAATGACTTC ATTCTGAGGG ATGGGACAGT CACCACAGAC TGGAAAGGCAC TCATCCAGGA
 6541 ATGGACCGTA CAGCAGCTTG GGAAGACATC CCAGCCTGTC CATGAGGGAGC AGTGTCTGT
 6601 CTCCCCAATTC TTCCACTGCCC AGGTCTCTT CTAGAATTG TTTGCCAGT GCCACAAGGT
 6661 CCTCCGCTCCA GCCACCTTTT ATGCCATGTG CCAGCCGAC AGTTGCCACC CGAACAAAGT
 6721 GTGTGAGGCG ATTGCCCTGT ATGCCACCT CTGTGGACC AAAGGGGTCT GTGTGGACTG
 6781 GAGGAGGGCC AATTTCTGTG CTATGTCATG TCCACCATCC CTGGTGTACA ACCACTGTGA
 6841 GCATGGCTGC CCTCGGCTCT GTGAAGCAA TACAAGCTCC TGTGGGGACC AACCCCTCGGA
 6901 AGGCTGCTTC TGCCCCCCAA ACCAAGTCAT GCTGGAAGGT AGCTGTGTCC CCGAGGAGGC
 6961 CTGTACCCAG TGCATCAGCG AGGATGGAGT CGGGCACCAG TTCCCTGGAAA CCTGGGTCCC
 7021 AGCCCACCAAG CTTGCCAGA TCTGCACGTG CCTCAGTGGG CGGAAGGTCA ACTGTACGTT
 7081 GCAGCCCTGC CCCACAGCCA AAGCTCCAC CTGTGGCCCG TGTGAAGTGG CCCGGCTCCG
 7141 CCAGAACGCA GTGCACTGCT GCCCCGAGTA CGAGTGTGTG TGTGACCTGG TGAGCTGTGA
 7201 CCTGGCCCCCG GTGCTCCTC GCAGAGATGG CCTCCAGATG ACCCTGACCA ATCCCTGGCGA
 7261 GTGCAAGACCC AACTCACT GTGCCCTGCAG GAAGGATGAA TGCAGACGGG AGTCCCCGCC
 7321 CTCTTGTCCC CGCACCGGA CGCCGGCCCT CGGAAGACT CAGTGTGTG ATGAGTATGA
 7381 GTGTGCATGC AACTGTGTCA ACTCCACGGT GAGCTGCCCG CTTGGGTACCG TGGCTCGGC
 7441 TGTCAACCAAC GACTGTGGCT GCACCCACAAAC TCTGCACCTT CCTGACAAGG TGTGTGTCCA
 7501 CCGAGGCACC ATCTACCCCTG TGGGCCAGTT CTGGGAGGAG GCCTGTGACG TGTGCACCTG
 7561 CACGGACTTG GAGGACTCTG TGATGGGCCT GCGTGTGGCC CAGTGTCTCC AGAACCCCTG
 7621 TGAGGACAAC TGCCTGTCA GCTTCACCTA TGTCTTCAAT GAAGGCGAGT GCTGTGGAAAG
 7681 GTGTCTGCCA TCTGCCCTGT AGGTGGTCAC TGGTTCAACCA CGGGGGAGC CCCAGTCTCA
 7741 CTGGAAGAAT GTGGCTCTC ACTGGGCCTC CCTGACAAAC CCCTGCTCA TCAATGAGTG
 7801 TGTCCGAGTG AAGGAAGAGG TCTTTGTCA ACAGAGGAAT GTCTCTGCC CCCAGCTGAA
 7861 TGTCCCCACC TGCCCCACGG GCTTCAGCT GAGCTGTAAAG ACCTCAGAGT GTTGTCCCAC
 7921 CTGTCACTGC GAGCCCCCTGG AGGCCTGCTT GCTCAATGGT ACCATCATTG GGCCGGGGAA
 7981 AAGTCTGATG ATTGATGTGT GTACACCTG CCGCTGCACC GTGCCGGTGG GAGTCATCTC
 8041 TGGATTCAAG CTGGAGGGCA GGAAGACAC CTTGTGAGGCA TGCCCCCTGG GTTATAAGGA
 8101 AGAGAAGAAC CAAGGTGAAT GCTGTGGAG ATGCTGTGCT ATAGCTTGCA CCATTCAGCT
 8161 AAGAGGAGGA CAGATCATGA CACTGAAGCG TGATGAGACT ATCCAGGATG GCTGTGACAG
 8221 TCACTTCTGC AAGGTCAATG AAAGAGGAGA GTACATCTGG GAGAAGAGAG TCACGGGTTG
 8281 CCCACCTTTC GATGAACACA AGTGTCTGCC TGAGGGAGGA AAAATCATGA AAATTCCAGG
 8341 CACCTGCTGT GACACATGTG AGGAGCCAGA ATGCAAGGAT ATCATTGCCA AGCTGCAGCG
 8401 TGTCAAAGTG GGAGACTGTA AGTCTGAAGA GGAAGTGGAC ATTCAATTACT GTGAGGGTAA
 8461 ATGTGCCAGC AAAGCCGTGT ACTCCATCCA CATGGAGGAT GTGCAGGACC AGTGCCTCTG
 8521 CTGCTCGCCC ACCCAGACGG AGCCCACGA GGTGGCCCTG CGCTGCACCA ATGGCTCCCT
 8581 CATCTACCAT GAGATCCTCA ATGCCATCGA ATGCAGGTGT TCCCCCAGGA AGTCAGCAA
 8641 GTGAGGCCAC TGCCTGGATG CTACTGTGCG CTGCCCTTACCGACCTCACT GGACTGGCCA
 8701 GAGTGTGCT CAGTCCCTCT CAGTCCCTCT CCTGCTCTGC TCTTGTGCTT CCTGATCCCCA
 8761 CAATAAAGGT CAATCTTCA CCTTGAAAAA AAAAAAAAAA AA

Human	MIPARFAGVILALALILPGTLCAEGTRGRSSTARCSLFSGSDFVNTPDGSMYSFAGYCSYL	60
Dog	-S-T-LVR-----K--TK--V----M----L-G--I----E----D----	
Human	LAGGCQKRSFSIIGDFQNGKRVSLSVYLGEFFDIHLFVNGETQGDQRVSMYASKGLYL	120
Dog	---D--EH-I-L--G---D-----ML--T-SI-----N----	
Human	ETEAGYYKLSGEAYGFVARIDSGGNFQVLLSDRYFNATCGLCGNFNIFAEDDFMTQEGTL	180
Dog	-A-----S-----N-----K-----	
Human	TSDPYDFANSWALSSGEQWCERASPPSSCNISSGEMQKGLWEQCQLLKSTSVFARCHPL	240
Dog	-----R-K-V----P--V--D-V-QV-----A-----	
Human	VDPEPFVALCEKTLCECAGGLECACPALLEYARTCAQEGMVLYGWTDHSACSPVCPAGME	300
Dog	-----R---T-VQ-M--P-AV-----A---Q-I-----V-R-A-----	
Human	YRQCVPSCARTCQLSHINEMCQERCVDGSCPEGQLLDEGLCVESTECPCVHSGKRYPPG	360
Dog	-KE-----T-----VK-V---Q-----H--G-A--S---A-Q-----	
Human	TSLSRDCNTCICPNSQWICSNEECPGECLVTGQSHFKSFDNRYFTFSGICQYLLARDQD	420
Dog	A--LQ--H-----L-----V-H-----Q-----	
Human	HSFSIVIETVQCADDRDAVCTRSVTVRPLGLHNSLVKLKHGAGVAWDCQDVQLPLKGDL	480
Dog	-T--V-----L-----H-----N-G--S-----I-I--Q-----	
Human	RIQHTVTASVRLSYGEDLQMDWDGRGRLLVKLSPVYAGKTCGLCGNYNGNQGDDFLTPSG	540
Dog	-----M-----S-V-----T-Y-A-----RG-----R-----V-A-----	
Human	LAEPRVEDFGNAWKLHGDCQDLQKQHSDPCALNPRMTRFSEEACAVLTSPTFEACHRAVS	600
Dog	-----L-----L-A-EN-----R--S---QA--A-----L---SK---P-----G	
Human	PLPYLRNCRYDVCSCSDGRECLCGALASYAAACAGRGRVVAWREPGRCELNCPKGQVYLO	660
Dog	-Q--VQ--L-----D---S-V-N---V-R---H-----F-A-S--Q-----	
Human	CGTPCNLTCSRSLSYPDEECNEACLEGCFCPGGLYMDERGLCVPKAQCPYCYYDGEIFQPED	720
Dog	-----M--L-----E-D---V--S--S-----L-----	
Human	IFSDKHTMCYCEDGFMHCTMSGVPGSLLPAVLSSPLSHRSKRSLSRPPMVKLVCPADN	780
Dog	-----T--GL-----NP-----RC-----	
Human	LRAEGLECTKTQNYDLECMMSGCVSGCLCPPGMVRHENRCAVALERCPCFHQGKEYAPGE	840
Dog	P-----A-----Q---T-----Q-----Q-----	
Human	TVKIGNTCVCRDRKWNTDHVCDATCSTIGMAHYLTFDGLKYLFPGEQYVLVQDYCGS	900
Dog	---D-----T-----A-----	
Human	NPGTFRILVGNKGCSHPVKCKRVTILVEGGEIELFDGEVNVKRPMKDETHFEVVESGR	960
Dog	---L-----E--Y-----K-----Q-----	
Human	YIILLLGKALSVVWDRHLSISVVLKQTYQEKCVCGLCGNFDGIQNNDLTSSNLQVEEDPVD	1020
Dog	-V-----HR-----T-R---Q-----F--S--I-----	
Human	FGNSWKVSSQCADTRKVPLDSSPATCHNNIMKQTMVDSSCRILTSDFQDCNKLVDPEPY	1080
Dog	-----NP-----K-----V-----I-----R-----F	

FIGURE 2A

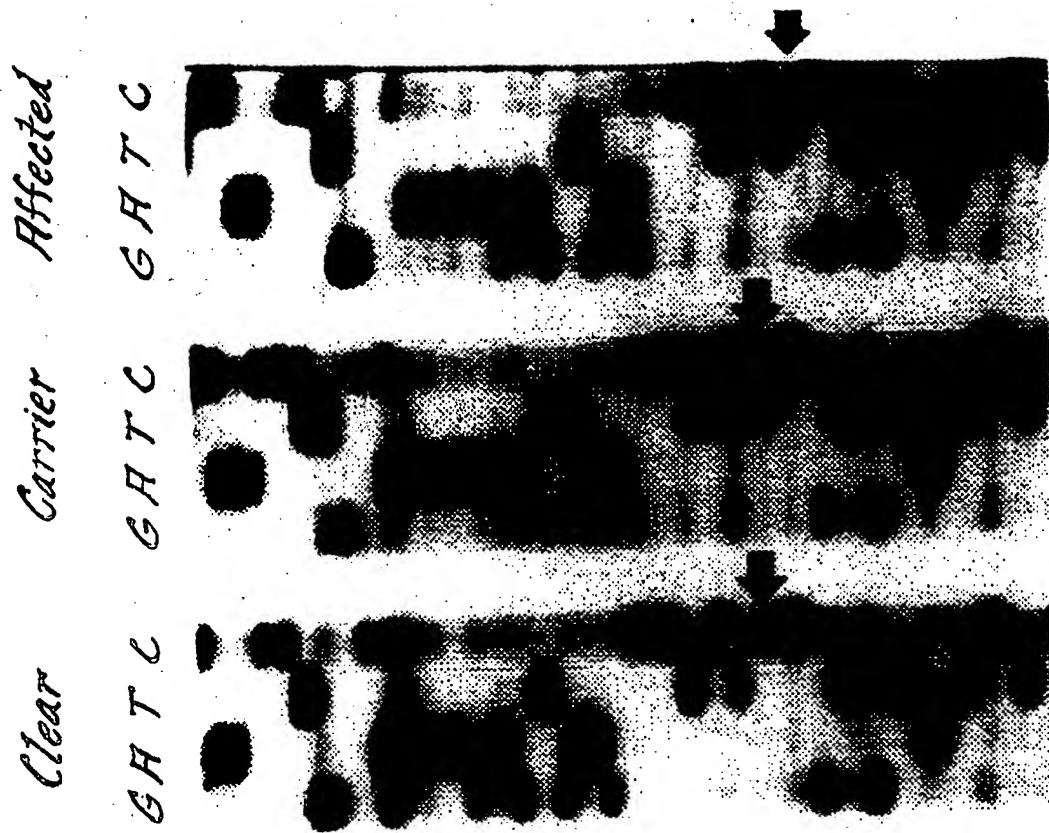
Human Dog	LDVCIYDTCSSESIGDCACFCDTIAAYAHVCAOHGVVTWRATLCPQSCEERNLRENGY --I-----T-----A-----F-----N-----H-----	1140
Human Dog	ECEWRYNSCAPACQVTQHQHPEPLACPVQCVEGCHAHCPPGKILDELLQTCVDPEDCPVCE -----PI-----I-----	1200
Human Dog	VAGRRFASGKKVTLNPSDPEHCQICHCDVNLTCACQEPGGLVVPPTDAPVSPPTLYVE ----L-P---II-----N-G-F-K-R-SV----G-IGS-S---	1260
Human Dog	DISEPPLHDFYCSRLLDLVFLLDGSSRLSEAEFEVLKAFVVDMMERLRISQKWVRVAVVE -T-----H-----K-----D-----V-----G-----H-H-----RI-----	1320
Human Dog	YHDGSHAYIGLKDRKRPSELRIASQVKYAGSQVASTSEVLKYTLFQIFSIDRPEASRI -----E-----T-----E-----G-----	1380
Human Dog	ALLLMASOEPORMSRNFVRYVQGLKKKVIVIPVGIGPHANLKQIRLIEKQAPENKAFVL -----S-LA--L-----S-----H-----F	1440
Human Dog	SSVDELEQQRDEIVSYLCDLAPEAPPPTLPPHMAQVTVGPGILLGVSTLGPKRNSMVLDVA -G-----R-----IN-----A-QH-P-----SE-----SP-----V	1500
Human Dog	FVLEGSDKIGEADFNRSYEFMEEVIQRMVGQDSIHVTVLQYSYMTVEYPFSEAOSKGD -----N-K-R-----R-----T-----E	1560
Human Dog	ILQRVREIRYQGGNRTNTGLALRYLSDHSFLVSQGDREQAPNLVYMTGNPASDEIKRLP V--Q--D--R-----Q---E---S-----V-----M-----	1620
Human Dog	GDIQVVPIGVGPNAVQELERIGWPNAPILODFETLPRAPDLVLQRCCSGEGLQIPTL -----H-----K-----H-----M-----	1680
Human Dog	SPAPDCSQPLDVILLLDGSSFPASYFDEMKSFAKAFISKANIGPRLTQSVLQYGSITT --T-----V-----I-----T-----R-----	1740
Human Dog	IDVPWNVVPEKAHLLSLVDVMQREGGPSQIGDALGFAVRYLTSEMHGARPAGSKAVVILV -----AY-V-----L-Q-----E-----S-----V-----V	1800
Human Dog	TDVSVDSDAAADAARSNRVTVPPIGIGDRYDAALQLRILAGPAGDSNVVKLQRIEDLPTM -----E-----SE---SS---KAG--M-R-----V	1860
Human Dog	VTLGNNSFLHKLCSGFVRICMDEDGNEKRPDVWTLPDQCHTVTCQPDGQTLKTHRVCND A-----F-----D-V-V-----L-----S-----	1920
Human Dog	RGLRPSCPNSQSPVKVEETCGCRWTCPCVCTGSSTRHIVTFDGQNFKLTGSCSYVLFQNK --P-----G-P-LR-----M-----	1980
Human Dog	EQDLEVLHNGACSPGARQGCMKSIEVKHSALSVELHSDMEVTVNGLVSVPYVGGNMEV -----Q-----KET-----DG-----QM-----I-----D-----	2040
Human Dog	NYYGAIMHEVRFNHLGHIFTFTPQNNEFQLQLSPKTFASKTYGLCGICDENGANDFMLRD ----T--Y-----R-----I-----	2100
Human Dog	GTVTTDWKTLVQEWTVQRPQOTCQPILEEQCLVPDSSHCVLPLFAECHKVLAATFY -----A-I-----QL-K-S--VH----P-SEFF-----SE-----	2160

FIGURE 2B

Human	AICQQDSCHQEQQVCEVIASYAHLCRTNGVCVDWRTPDFCAMSCPPSLVYNHCEHGCPRHC	2220
Dog	-M---P---PKK---A---L---K---RAN-----L-	
Human	DGNVSSCGDHPSEGCFCPPDKVMLEGSCVPEEACTQCIGEDGVQHQFLEAWVPDFQPCQI	2280
Dog	E---T---Q-----NQ-----S---R---T---A-----	
Human	CTCLSGRKVNCTTQPCPTAKAPTCGLCEVARLRQNADQCCPEYECVCDPVSCDLPPVPHC	2340
Dog	-----L-----P-----V-----L-----P-	
Human	ERGLQPTLTNPGECRPNFTCACRKEECKRVSPPSCPPHRLPTLRKTQCCDEYEACNCVN	2400
Dog	-D---M-----D---R-E-----T-A-----	
Human	STVSCPPLGYLASTATNDGCCTTCLPDKVCVHRSTIYPVGQFWEEGCDVCTCTDMEDAV	2460
Dog	-----AV-----F-----G-----A-----L---S-	
Human	MGLRVAQCSQKPCEDSCRSGFTYVLHEGECCGRCLPSACEVVTGS	2520
Dog	-----N-L-----[REDACTED]-----A---H---N---H	
Human	WASPENPCLINECVRVKEEVFIQQRJNVSCPQLEVPCPSGFQLSCKTSACCPSCRERME	2580
Dog	-----D-----V-----N---T---T-----E---T-H---PL-	
Human	ACMNLNGTIVIGPGKTVMDVCTTCRGMVQGVVISGFKLECRKTCNPCPLGYKEENNTGEC	2640
Dog	--L---I---SL-----T-P-----G-----EA-----K-Q---	
Human	CGRCLPTACTIQLRGQIMTLKRDETLQDGCDTHFKVNERGEYFWEKRVITGCPPFDEHK	2700
Dog	-----I-----I-----S-----I-----	
Human	CLAEGGKIMKIPGTCCDTCEEPECNDITARLQYVKVGSKSEVEVDIHYCOGKCASKAMY	2760
Dog	-----K---I---R---D---E-----E-----V-	
Human	SIDINDVQDQCSCCSPTRTEPMQVALHCTNGSVVYHEVLNAMECKCSPRKCSK	2813
Dog	--HME-----Q-----R-----LI---I---I---R-----	

FIGURE 2C

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1. 2.

3. 4.

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exon 4 AAATGACAAAAGAGTGAGCCGGTC*

AGGGGGTTTCCAAAATGACAAAAGAGTGAGCCTCTCCGTGTATCTCGGAGAATTTTCGA
G G F Q N D K R V S L S V Y L G E F F D

CATTCATTTGTTGTCAATGGTACCATGCTGCAGGGACCCAAAGGTAAAGTCAGAAGCCC
I H L F V N G T M L Q G T Q R

GAATGTTCAGGTTAATATGGACCCCTGGGATCACTTGCAACCCCCTGTTTTTCAGAT

GAGGGAGCCGGGGCCCAGAGACAGGAAGTAAATGTGCCAGGGAAAGTGAGTGGCAGGAC

TGGGTGAAAGCCCCATATCCCGACTCCTGGTCAAGGAGACTTGCACCAAGGTCCCAGCC
3'-GGGCTGGCGACCAGTTCCTCTGAA-5'

CTGGAGCATGGGTTGGGTTGGAAGGTGGAGGGACATGGAGGAAATGCATGAGAACAC

exon 5
GCTTCCTGAGCTCCTCCTGTCCCACCAGCATCTCCATGCCCTACGCCCTCCAATGGGC
I S M P Y A S N G

FIGURE 4

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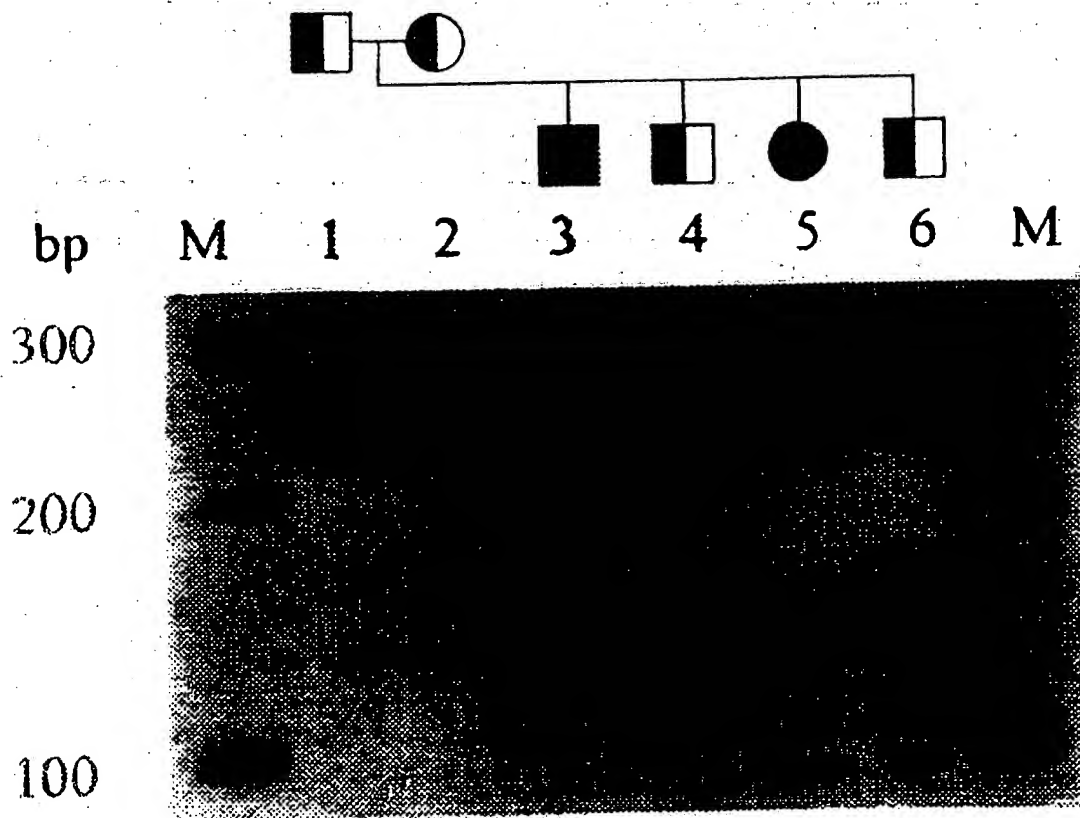


Fig. 5.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US97/12606

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) :C12Q 1/68; C12P 19/34; C07H 21/02, 21/04

US CL :435/6, 91.2; 536/23.1, 24.3, 24.33

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 91.2; 536/23.1, 24.3, 24.33

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Extra Sheet.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y --- A	SHIBUYA, H. et al. A polymorphic (AGGAAT) _n tandem repeat in an intron of the canine von Willebrand factor gene. Animal Genetics. April 1994, Volume 25, Number 2, page 122, see entire document.	15-22, 24-26, 28, 31 ----- 1-14, 23, 27, 29

Further documents are listed in the continuation of Box C. See patent family annex.

•	Special categories of cited documents:	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
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Date of the actual completion of the international search
28 AUGUST 1997

Date of mailing of the international search report

14 NOV 1997

Name and mailing address of the ISA/US
Commissioner of Patents and Trademarks
Box PCT
Washington, D.C. 20231
Facsimile No. (703) 305-3230

Authorized officer
DIANNE REES
Telephone No. (703) 308-0196

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US97/12606

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, BIOSIS, BIOTECHABS, BIOTECHDS, CABA, DGENE, DRUGU, EMBASE, MEDLINE, EUROPATFULL,
JAPIO, WPIDS, USPATFULL, GENBANK

search terms: von Willebrand, sequence, clone, cloning, probes, primers, hybridization, detection, nucleic acids, mutations, canine, dogs, Scottish terriers, primers in Figure 4.

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